



## METHODOLOGY

Orphanet provides a comprehensive inventory of rare diseases in Europe, published biannually as a list. Rare diseases registered in Orphanet are defined according to two scopes:

- Every entity is defined by its clinical homogeneity, regardless of its etiology or the number of causing genes identified;
- The rarity is defined according to the European legislation defining a prevalence threshold of not more than 5 affected persons per 10'000 (Regulation (EC) N°141/2000 of the European Parliament and of the Council of 16 December 1999 on orphan medicinal products, [http://ec.europa.eu/health/files/eudralex/vol-1/reg\\_2000\\_141/reg\\_2000\\_141\\_en.pdf](http://ec.europa.eu/health/files/eudralex/vol-1/reg_2000_141/reg_2000_141_en.pdf)).

Registered rare diseases have been described in the international scientific literature (peer-reviewed articles) with at least two cases confirming that the clinical signs are not associated fortuitously. However, some diseases are registered although only one case has been reported in order to reproduce the comprehensiveness of a specific classification (notably within the classification of inborn errors of metabolism).

Rare diseases are registered with a preferred name and as many synonyms as necessary. A unique identifier, the ORPHA number, is randomly attributed by the database to each disease. This number is never re-used, so it is stable in time.

ORPHA number of rare diseases registered in the past may be absent from the current inventory. This is due to:

- Obsolescence of entries (e.g. duplicated entities, diseases that are not rare anymore);
- Deprecation of entities when an entity no longer exists per se but has been recognised as being another entity. In this case, information regarding the deprecated entity is moved and the users are redirected to the target entry.

### Data collection

As new scientific knowledge arises, the Orphanet rare diseases inventory is updated through the regular addition/update of diseases via two non-exclusive sources: documented sources and/or expert advice.

The scientific knowledge is monitored through:

- A bi-monthly analysis of a defined set of international peer-reviewed scientific journals covering the diversity of medical specialities represented in Orphanet;
- A monthly Medline search algorithm: (nosology[Title] OR classification[Title] OR nomenclature[Title] OR terminology[Title]) AND (rare disease\* OR syndrome\* OR disorder\*);
- Specific Medline queries according to requests from experts, users of the database or needs arising from services newly registered in Orphanet (e.g. diagnostic test, expert centre, patient organisation).

Update of the inventory of rare diseases is assessed monthly by a medical and scientific committee within Orphanet and further validated by consulted experts.

The Orphanet rare disease nomenclature is produced in English and is translated into other languages. A medical validation of the translations is carried out.

### Data presentation

Preferred names and synonyms of diseases are listed by alphabetical order with their ORPHA number.

Deprecated entities are listed with the ORPHA number to be used preceded by the sign “→ “. A table in annex lists the name of the rare disease and its ORPHA number to be used instead of the deprecated entries.

Obsolete entries are not listed here. In the case of duplicates, the nomenclature of the obsolete entry has been added to the rare disease listed here.

## Rare diseases listed in alphabetical order

ORPHA number	Disease name	ORPHA number	Disease name
289157	1-alpha-hydroxylase deficiency	352328	3-methylglutaconic aciduria with deafness-encephalopathy-Leigh-like syndrome
431361	2,4-dienoyl-CoA reductase deficiency	1772	45,X0/46,XY mixed gonadal dysgenesis
976	2,8-dihydroxyadenine urolithiasis	243	46,XX complete gonadal dysgenesis
79154	2-aminoadipic 2-oxoadipic aciduria	2973	46,XX disorder of sex development-anorectal anomalies syndrome
391417	2-methyl-3-hydroxybutyric aciduria	67046	3-methylglutaconyl-CoA hydratase deficiency
391428	2-methyl-3-hydroxybutyric aciduria, classic type	134	3-oxothiolase deficiency
391428	2-methyl-3-hydroxybutyric aciduria, infantile type	79351	3-phosphoglycerate dehydrogenase deficiency, infantile/juvenile form
391457	2-methyl-3-hydroxybutyric aciduria, neonatal type	79350	3-phosphoserine phosphatase deficiency
391417	2-methyl-3-hydroxybutyryl-CoA dehydrogenase deficiency	869	3A syndrome
391428	2-methyl-3-hydroxybutyryl-CoA dehydrogenase deficiency, classic type	7	3C syndrome
391428	2-methyl-3-hydroxybutyryl-CoA dehydrogenase deficiency, infantile type	2616	3M syndrome
391457	2-methyl-3-hydroxybutyryl-CoA dehydrogenase deficiency, neonatal type	293843	3MC syndrome
79095	2-methylacyl-CoA racemase deficiency	→293843	3MC1 syndrome
79157	2-methylbutyric aciduria	→293843	3MC2 syndrome
79157	2-methylbutyryl-CoA dehydrogenase deficiency	→293843	3MC3 syndrome
255182	2-oxoglutarate complex deficiency	67046	3MG-CoA hydratase deficiency
869	2A syndrome	2118	4-HPPD deficiency
2616	3-M syndrome	2118	4-alpha-hydroxyphenylpyruvate hydroxylase deficiency
2671	3-Phosphoglycerate dehydrogenase deficiency, neonatal form	22	4-hydroxybutyric aciduria
79301	3-beta-hydroxy-delta-5-C27-steroid oxidoreductase deficiency	2118	4-hydroxyphenylpyruvic acid dioxygenase deficiency
20	3-hydroxy-3-methylglutaric aciduria	869	4A syndrome
20	3-hydroxy-3-methylglutaryl-CoA lyase deficiency	88637	4H syndrome
35701	3-hydroxy-3-methylglutaryl-CoA synthase deficiency	250977	5-amino-4-imidazole carboxamide ribosiduria
939	3-hydroxyisobutyric aciduria	217064	5-fluorouracil intoxication
134	3-ketothiolase deficiency	217064	5-fluorouracil poisoning
6	3-methylcrotonyl-CoA carboxylase deficiency	33572	5-oxoprolinase deficiency
6	3-methylcrotonylglycinuria	99135	6-phosphogluconate dehydrogenase deficiency
67046	3-methylglutaconic aciduria type 1	13	6-pyruvoyl-tetrahydropterin synthase deficiency
111	3-methylglutaconic aciduria type 2	818	7-dehydrocholesterol reductase deficiency
67047	3-methylglutaconic aciduria type 3	168588	11-beta-hydroxysteroid dehydrogenase deficiency type 1
67048	3-methylglutaconic aciduria type 4	320	11-beta-hydroxysteroid dehydrogenase deficiency type 2
66634	3-methylglutaconic aciduria type 5	752	17-beta-hydroxysteroid dehydrogenase 3 deficiency
445038	3-methylglutaconic aciduria type 7	752	17-ketoreductase deficiency
		752	17-ketosteroidreductase deficiency
		99763	18-hydroxylase deficiency
		99763	18-oxidase deficiency
		881	45,X syndrome
		881	45,X/46,XX syndrome
		1772	45,X/46,XY MGD
		1772	45,X/46,XY mixed gonadal dysgenesis
		1772	45,X0/46,XY MGD
		1772	45,X0/46,XY mixed gonadal dysgenesis
		243	46,XX complete gonadal dysgenesis
		2973	46,XX disorder of sex development-anorectal anomalies syndrome
		2975	46,XX disorder of sex development-skeletal anomalies syndrome
		243	46,XX gonadal dysgenesis
		243	46,XX ovarian dysgenesis
		444048	46,XX ovarian dysgenesis-short stature syndrome
		2138	46,XX ovotesticular DSD
		2138	46,XX ovotesticular disorder of sex development
		243	46,XX pure gonadal dysgenesis
		393	46,XX testicular DSD
		393	46,XX testicular disorder of sex development
		199310	46,XX/46,XY chimerism
		242	46,XY CGD
		753	46,XY DSD due to 5-alpha-reductase 2 deficiency
		755	46,XY DSD due to LH resistance or LHB deficiency
		325448	46,XY DSD due to LHB deficiency
		96265	46,XY DSD due to complete LH receptor inactivation
		96265	46,XY DSD due to complete LH resistance
		96265	46,XY DSD due to complete luteinizing hormone receptor inactivation
		96265	46,XY DSD due to complete luteinizing hormone resistance
		755	46,XY DSD due to luteinizing hormone resistance or luteinizing hormone beta subunit deficiency
		325448	46,XY DSD due to luteinizing hormone subunit beta deficiency
		96266	46,XY DSD due to partial LH receptor inactivation
		96266	46,XY DSD due to partial LH resistance
		96266	46,XY DSD due to partial luteinizing hormone resistance
		251510	46,XY PGD
		242	46,XY complete gonadal dysgenesis
		96266	46,XY disorder of sex development due to partial LH receptor inactivation
		96266	46,XY disorder of sex development due to partial LH resistance
		96266	46,XY disorder of sex development due to partial luteinizing hormone resistance

→ This disease is deprecated and has been moved to another (see annex). The indicated Orphan number should now be used.

ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
752	46,XY disorder of sex development due to 17-beta-hydroxysteroid dehydrogenase 3 deficiency	293948	1p21.3 microdeletion syndrome	459074	7q36.3 microduplication syndrome
		401986	1p31p32 microdeletion syndrome	96092	8p inverted duplication/deletion syndrome
		456298	1p35.2 microdeletion syndrome		
753	46,XY disorder of sex development due to 5-alpha-reductase 2 deficiency	1606	1p36 deletion syndrome	168953	8p11 myeloproliferative syndrome
		250989	1q21.1 microdeletion syndrome	251066	8p11.2 deletion syndrome
		250994	1q21.1 microduplication syndrome	251076	8p23.1 duplication syndrome
755	46,XY disorder of sex development due to LH resistance or LHB deficiency	250999	1q41-q42 microdeletion syndrome	251071	8p23.1 microdeletion syndrome
		250999	1q41q42 microdeletion syndrome	228399	8q12 microduplication syndrome
		238769	1q44 microdeletion syndrome	2496	8q13 microdeletion syndrome
325448	46,XY disorder of sex development due to LHB deficiency	363680	2p13.2 microdeletion syndrome	284160	8q21.11 microdeletion syndrome
96265	46,XY disorder of sex development due to complete LH receptor inactivation	261349	2p15-p16.1 microdeletion syndrome	178303	8q22.1 microdeletion syndrome
		261349	2p15p16.1 microdeletion syndrome	261112	9p deletion syndrome
		163693	2p21 deletion syndrome	261112	9p- syndrome
96265	46,XY disorder of sex development due to complete LH resistance	163693	2p21 microdeletion syndrome	324313	9p13 microdeletion syndrome
		369881	2p21 microdeletion syndrome without cystinuria	96147	9q subtelomeric deletion syndrome
96265	46,XY disorder of sex development due to complete luteinizing hormone receptor inactivation	228402	2q23.1 microdeletion syndrome	352665	9q21 microdeletion syndrome
		313947	2q23.1 microduplication syndrome	401923	9q31.1q31.3 microdeletion syndrome
		1617	2q24 microdeletion syndrome		
96265	46,XY disorder of sex development due to complete luteinizing hormone resistance	251014	2q31.1 microdeletion syndrome	284169	10p12p11 microdeletion syndrome
		294026	2q31.1 microduplication syndrome	276413	10q22.3q23.3 microdeletion syndrome
		251019	2q32-q33 microdeletion syndrome		
90796	46,XY disorder of sex development due to isolated 17,20-lyase deficiency	251019	2q32q33 microdeletion syndrome	276422	10q22.3q23.3 microduplication syndrome
		251028	2q33.1 microdeletion syndrome		
		1001	2q37 microdeletion syndrome		
755	46,XY disorder of sex development due to luteinizing hormone resistance or luteinizing hormone beta subunit deficiency	1620	3p- syndrome	1307	10q24 microduplication syndrome
		435638	3p25.3 microdeletion syndrome	52022	11p11.2 deletion
		65286	3q subtelomere deletion syndrome	300305	11p15.4 microduplication syndrome
325448	46,XY disorder of sex development due to luteinizing hormone subunit beta deficiency	65286	3qter deletion	444002	11q22.2-q22.3 deletion syndrome
		1621	3q13 microdeletion syndrome	444002	11q22.2q22.3 microdeletion syndrome
		96095	3q26 microduplication syndrome	313884	12p12.1 microdeletion syndrome
443087	46,XY disorder of sex development due to testicular 17,20-desmolase deficiency	356947	3q26-q27 microdeletion syndrome	280325	12p13.33 microdeletion syndrome
		356947	3q26q27 microdeletion syndrome	94063	12q14 microdeletion syndrome
		397695	3q27.3 microdeletion syndrome	289513	12q15q21.1 microdeletion syndrome
168558	46,XY disorder of sex development-adrenal insufficiency due to CYP11A1 deficiency syndrome	65286	3q29 microdeletion syndrome	412035	13q12.3 microdeletion syndrome
168563	46,XY gonadal dysgenesis-motor and sensory neuropathy syndrome	251038	3q29 microduplication	1590	13q32 deletion
		280	4p- syndrome	261120	14q11.2 microdeletion syndrome
		96072	4p16.3 microduplication syndrome	261229	14q11.2 microduplication syndrome
325345	46,XY ovotesticular DSD	238750	4q21 microdeletion syndrome	261144	14q12 microdeletion syndrome
325345	46,XY ovotesticular disorder of sex development	329802	5p13 microduplication syndrome	→3157	14q22 microdeletion syndrome
		86841	5q- syndrome	264200	14q22-q23 microdeletion syndrome
		228384	5q14.3 microdeletion syndrome	264200	14q22q23 microdeletion syndrome
251510	46,XY partial gonadal dysgenesis	436003	5q23 microdeletion syndrome	401935	14q24.1q24.3 microdeletion syndrome
251510	46,XY partial testicular dysgenesis	314655	5q31.3 microdeletion syndrome	314585	15q overgrowth syndrome
242	46,XY pure gonadal dysgenesis	228415	5q35 microduplication syndrome		
3375	47,XXX syndrome	96125	6p subtelomeric deletion syndrome	238446	15q11-q13 duplication syndrome
8	47,XYY syndrome	251046	6p22 microdeletion syndrome	238446	15q11-q13 microduplication syndrome
9	48,XXXX syndrome	96125	6p25 microdeletion syndrome		
96263	48,XXXY syndrome	75857	6q terminal deletion syndrome	261183	15q11.2 microdeletion syndrome
10	48,XXYY syndrome	171829	6q16 deletion syndrome	238446	15q11q13 duplication syndrome
99329	48,XYYY syndrome	251056	6q25 microdeletion syndrome	238446	15q11q13 microduplication syndrome
11	49,XXXXX syndrome	314034	7p22.1 microduplication syndrome		
96264	49,XXXXY syndrome	96121	7q11.23 microduplication syndrome	199318	15q13.3 microdeletion syndrome
261534	49,XXYY syndrome	251061	7q31 microdeletion syndrome		
99330	49,YYYY syndrome				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
261190	15q14 microdeletion syndrome	268261	21q22.13q22.2 microdeletion syndrome	69739	ABSD
94065	15q24 microdeletion syndrome			2310	Absence deformity of leg-cataract syndrome
1596	15q26 deletion syndrome	567	22q11DS	99112	Absence of brachiocephalic vein
363992	15q26.3 microdeletion syndrome	567	22q11.2 deletion syndrome	1658	Absence of dermatoglyphics-congenital milia syndrome
261211	16p11.2-p12.2 microdeletion syndrome	1727	22q11.2 microduplication syndrome		
261211	16p11.2p12.2 microdeletion syndrome	48652	22q13 deletion	1658	Absence of fingerprints-congenital milia syndrome
		85445	AA amyloidosis		
261204	16p11.2p12.2 microduplication syndrome	869	AAA syndrome	99112	Absence of innominate vein
		35708	AADC deficiency	101206	Absence of pulmonary valve-Fallot tetralogy-absence of ductus arteriosus syndrome
485405	16p12.1p12.3 triplication syndrome	91385	AAE		
261236	16p13.11 microdeletion syndrome	100055	AAE 2	99048	Absence of pulmonary valve-ventricular septal defect-persistent ductus arteriosus syndrome
261243	16p13.11 microduplication syndrome	100055	AAE II		
96078	16p13.3 microduplication syndrome	1414	Aagenaes syndrome	980	Absence of the pulmonary artery
		284460	AAOR		
352629	16q24.1 microdeletion syndrome	93560	AApoAI amyloidosis	99114	Absence of the superior caval vein
261250	16q24.3 microdeletion syndrome	238269	AApoAII amyloidosis	99114	Absence of the superior vena cava
819	17p11.2 microdeletion syndrome	439232	AApoAIV amyloidosis	99114	Absence of the SVC
1713	17p11.2 microduplication syndrome	915	Aarskog syndrome	1493	Absent corpus callosum-cataract-immunodeficiency syndrome
477817	17p11.2p12 microduplication syndrome	1974	Aarskog-like syndrome		
217385	17p13.3 duplication syndrome	3163	Aarskog-Ose-Pande syndrome	85201	Absent patellae-scrotal hypoplasia-renal anomalies-facial dysmorphism-intellectual disability syndrome
		915	Aarskog-Scott syndrome		
217385	17p13.3 microduplication syndrome	124	Aase syndrome	3016	Absent radius-anogenital anomalies syndrome
97685	17q11 microdeletion syndrome	916	Aase-Smith I syndrome		
139474	17q11.2 microduplication syndrome	124	Aase-Smith II syndrome	2951	Absent thumb-short stature-immunodeficiency syndrome
261265	17q12 microdeletion syndrome	916	Aase-Smith syndrome		
261272	17q12 microduplication syndrome	69663	ABCB4 gene mutation-associated cholelithiasis	988	Absent tibia-polydactyly syndrome
363958	17q21.31 microdeletion syndrome	→897	ABCD syndrome	3328	Absent tibia-polydactyly-arachnoid cyst syndrome
217340	17q21.31 microduplication syndrome	2970	Abdominal muscle deficiency syndrome	99901	ACAD9 deficiency
261279	17q23.1-q23.2 microdeletion syndrome	800	Aberfeld syndrome	42	ACADM deficiency
261279	17q23.1q23.2 microdeletion syndrome	85446	ABeta2Mwt amyloidosis	26792	ACADS deficiency
		324723	ABeta amyloidosis, Arctic type	945	Acalvaria
1598	18p- syndrome	100006	ABeta amyloidosis, Dutch type	67043	Acanthamoeba keratitis
1600	18q- syndrome	324718	ABeta amyloidosis, Flemish type	79468	Acanthokeratolytic verrucous nevus
254346	19p13.12 microdeletion syndrome	324708	ABeta amyloidosis, Iowa type	300504	Acanthoma of the nail matrix
357001	19p13.13 microdeletion syndrome	324713	ABeta amyloidosis, Italian type	90301	Acanthosis nigricans-insulin resistance-muscle cramps-acral enlargement syndrome
447980	19p13.3 microduplication syndrome	324703	ABeta amyloidosis, Piedmont type		
217346	19q13.11 microdeletion syndrome	324718	ABetaA21G amyloidosis	926	Acatalasemia
313781	20p subtelomeric deletion syndrome	324718	ABetaA21G-related amyloidosis	2508	ACC-abnormal genitalia syndrome
261295	20p12.3 microdeletion syndrome	324708	ABetaD23N amyloidosis	561	Accelerated skeletal maturation-facial dysmorphism-failure to thrive syndrome
313781	20p13 microdeletion syndrome	324723	ABetaE22G amyloidosis		
444051	20q11.2 microdeletion syndrome	324713	ABetaE22K amyloidosis	180182	Accessory breasts
363659	20q11.2 microduplication syndrome	100006	ABetaE22Q amyloidosis		
261311	20q13.33 microdeletion syndrome	324703	ABetaL34V amyloidosis	99061	Accessory mitral valve tissue
574	21q deletion syndrome	324703	ABetaL34V-related amyloidosis	141096	Accessory nostril
574	21q- syndrome	14	Abetalipoproteinemia	674	Accessory pancreas
261323	21q22.11-q22.12 microdeletion syndrome	920	Ablepharon macrostomia syndrome	95462	Accessory tricuspid valve tissue
		99089	Abnormal number of coronary ostia	210122	ACDMPV
261323	21q22.11q22.12 microdeletion syndrome	99050	Abnormal origin of right or left pulmonary artery from the aorta	48818	Aceruloplasminemia
268261	21q22.13-q22.2 microdeletion syndrome	1164	ABPA	464458	Acetaminophen poisoning
		97345	ABri amyloidosis		
		921	Abruzzo-Erickson syndrome		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
99736	Acetazolamide-responsive congenital myotonia	100056	Acquired angioneurotic edema type 1	93585	Acquired thrombotic thrombocytopenic purpura
99736	Acetazolamide-responsive myotonia	100055	Acquired angioneurotic edema type 2	93585	Acquired TTP
2008	ACFS			99147	Acquired von Willebrand disease
930	Achalasia cardia	91385	Acquired bradykinine-induced angioedema	99147	Acquired von Willebrand syndrome
869	Achalasia-addisonianism-alacrima syndrome	91385	Acquired C1 inhibitor deficiency	263534	Acral deciduous skin
→869	Achalasia-alacrimia syndrome	95626	Acquired CDI	97360	Acral dysostosis with facial and genital abnormalities
929	Achalasia-microcephaly syndrome	95626	Acquired central diabetes insipidus	158673	Acral dystrophic epidermolysis bullosa
294983	Acheiria	454700	Acquired Creutzfeldt-Jakob disease		
295103	Acheiria, bilateral	228285	Acquired cutis laxa	263534	Acral peeling skin syndrome
295101	Acheiria, unilateral	404514	Acquired cystic disease-associated renal cell carcinoma	90396	Acral persistent papular mucinosis
931	Acheiropodia	46487	Acquired epidermolysis bullosa	263534	Acral PSS
931	Acheiropody	98818	Acquired epileptic aphasia	281127	Acral self-healing collodion baby
49382	ACHM	91136	Acquired Fanconi syndrome secondary to monoclonal gammopathy	281127	Acral SHCB
932	Achondrogenesis	91136	Acquired Fanconi syndrome secondary to monoclonal gammopathy	945	Acrania
93299	Achondrogenesis type 1A			978	Acro-dermato-ungual-lacrima-tooth syndrome
93298	Achondrogenesis type 1B	79086	Acquired generalized lipodystrophy	958	Acro-renal-mandibular syndrome
93296	Achondrogenesis type 2	228247	Acquired Gronblad-Strandberg-Touraine syndrome	959	Acro-renal-ocular syndrome
93299	Achondrogenesis, Houston-Harris type	231401	Acquired HbH disease	36	Acrocallosal syndrome
93296	Achondrogenesis, Langer-Saldino type	231401	Acquired hemoglobin H disease	63446	Acrocapitofemoral dysplasia
93298	Achondrogenesis, Parenti-Fraccaro type	158057	Acquired hemophagocytic lymphohistiocytosis associated with malignant disease	2008	Acrocardiofacial syndrome
15	Achondroplasia	73274	Acquired hemophilia	221054	Acrocephalopolydactylous dysplasia
935	Achondroplasia-SCID syndrome	2221	Acquired hypertrichosis lanuginosa	221054	Acrocephalopolydactyly
935	Achondroplasia-severe combined immunodeficiency syndrome	26348	Acquired hypoprothrombinemia	65759	Acrocephalopolysyndactyly type 2
935	Achondroplasia-Swiss type agammaglobulinemia syndrome	454	Acquired ichthyosis	65798	Acrocephalopolysyndactyly type 4
49382	Achromatopsia	75564	Acquired idiopathic sideroblastic anemia	87	Acrocephalosyndactyly type 1
355	Acid beta-glucosidase deficiency	37559	Acquired kinky hair syndrome	794	Acrocephalosyndactyly type 3
333	Acid ceramidase deficiency	79086	Acquired lipatrophic diabetes	710	Acrocephalosyndactyly type 5
35121	Acid phosphatase deficiency	464453	Acquired methemoglobinemia	63440	Acrocephaly
424046	Acinar cell carcinoma of pancreas	91136	Acquired monoclonal Ig light chain-associated Fanconi syndrome	949	Acrocraniofacial dysostosis
40366	Acitretin/etretinate embryopathy	91136	Acquired monoclonal immunoglobulin light chain-associated Fanconi syndrome	955	Acrodentoosteodysplasia
79099	Ackerman dermatitis syndrome	589	Acquired myasthenia	163931	Acrodermatitis continua suppurativa of Hallopeau
2561	Ackerman syndrome	95626	Acquired neurogenic diabetes insipidus	37	Acrodermatitis enteropathica
51890	ACNES	84142	Acquired neuromyotonia	978	Acrodermatoungualacrima-tooth syndrome
43115	Aconitase deficiency	91385	Acquired non histamine-induced angioedema	950	Acrodysostosis
252175	Acoustic neurilemoma	314697	Acquired porencephaly	280651	Acrodysostosis with multiple hormone resistance
252175	Acoustic neurinoma	729	Acquired primary erythrocytosis	950	Acrodysplasia
252175	Acoustic neuroma	26348	Acquired prothrombin deficiency	2956	Acrodysplasia scoliosis
65759	ACPS2	228247	Acquired pseudoxanthoma elasticum	1786	Acrofacial dysostosis, Catania type
65798	ACPS4	49566	Acquired purpura fulminans	246	Acrofacial dysostosis, Genee-Wiedmann type
306431	Acquired adult-onset immunodeficiency	228247	Acquired PXE	64542	Acrofacial dysostosis, Kennedy-Teebi type
90065	Acquired aneurysmal subarachnoid hemorrhage	206575	Acquired rippling muscle disease	1787	Acrofacial dysostosis, Palagonia type
91385	Acquired angioedema	485275	Acquired schizencephaly	1788	Acrofacial dysostosis, Rodríguez type
100056	Acquired angioedema type 1			952	Acrofacial dysostosis, Weyers type
100055	Acquired angioedema type 2			1784	Acrofrontofacionasal dysostosis
91385	Acquired angioneurotic edema				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2211	Acrofrontofacionasal dysostosis type 2	397596	Activated PI3K-delta syndrome	178320	Acute lung injury
2211	Acrofrontofacionasal syndrome type 2	101089	Activation-induced cytidine deaminase deficiency	518	Acute megakaryoblastic leukemia
		73423	Acute ackee fruit intoxication	99887	Acute megakaryoblastic leukemia in Down syndrome
2500	Acrogeria	95409	Acute adrenal failure	329469	Acute megakaryoblastic leukemia without Down syndrome
2500	Acrogeria, Gottron type	95409	Acute adrenal insufficiency		
38	Acrokeratoelastoidosis of Costa	95409	Acute adrenocortical insufficiency	518	Acute megakaryocytic leukemia
166113	Acrokeratosis of Bazex	99870	Acute and disseminated Langerhans cell histiocytosis	514	Acute monoblastic leukemia
166113	Acrokeratosis paraneoplastica			514	Acute monocytic leukemia
166113	Acrokeratosis paraneoplastica of Bazex	284460	Acute annular outer retinopathy	98917	Acute motor and sensory axonal neuropathy
		86849	Acute basophilic leukemia		
79151	Acrokeratosis verruciformis of Hopf	98837	Acute biphenotypic leukemia	98918	Acute motor axonal neuropathy
965	Acromegaloid facial appearance syndrome	2901	Acute brachial plexus neuritis	98917	Acute motor-sensory axonal GBS
963	Acromegaly	83597	Acute disseminated encephalitis	98917	Acute motor-sensory axonal Guillain-Barré syndrome
→2796	Acromegaly-cutis verticis gyrata-corneal leukoma syndrome	83597	Acute disseminated encephalomyelitis	228157	Acute multiple sclerosis, Marburg type
39	Acromelanosis	163703	Acute encephalitis with refractory repetitive partial seizures	228157	Acute multiple sclerosis, Marburg variant
1827	Acromelic frontonasal dysostosis	363549	Acute encephalopathy with biphasic seizures and late reduced diffusion	520	Acute myeloblastic leukemia 3
1827	Acromelic frontonasal dysplasia				
968	Acromesomelic dwarfism	279888	Acute endophthalmitis	98833	Acute myeloblastic leukemia M1
2098	Acromesomelic dysplasia, Grebe type	318	Acute erythroid leukemia	98834	Acute myeloblastic leukemia M2
		243367	Acute fatty liver of pregnancy	98834	Acute myeloblastic leukemia with maturation
968	Acromesomelic dysplasia, Hunter-Thompson type	3243	Acute febrile neutrophilic dermatosis	98833	Acute myeloblastic leukemia without maturation
40	Acromesomelic dysplasia, Maroteaux type	293173	Acute generalized exanthematous pustulosis	86843	Acute myelodysplasia with myelofibrosis
2500	Acrometageria	99920	Acute graft versus host disease	86843	Acute myelofibrosis
969	Acromicric dysplasia	90062	Acute hepatic failure	102379	Acute myeloid leukemia and myelodysplastic syndromes related to alkylating agent
955	Acroosteolysis dominant type	98916	Acute idiopathic demyelinating polyneuropathy		
955	Acroosteolysis with osteoporosis and changes in skull and mandible	363549	Acute infantile encephalopathy predominantly affecting the frontal lobes	164726	Acute myeloid leukemia and myelodysplastic syndromes related to radiation
363665	Acroosteolysis-keloid-like lesions-premature aging syndrome				
2980	Acrotoocular syndrome	217371	Acute infantile liver failure due to synthesis defect of mitochondrial DNA-encoded proteins	102381	Acute myeloid leukemia and myelodysplastic syndromes related to topoisomerase type 2 inhibitor
85203	Acropectoral syndrome				
956	Acropectorrenal dysplasia	217371	Acute infantile liver failure due to synthesis defect of mtDNA-encoded proteins	318	Acute myeloid leukemia M6
957	Acropectorovertebral dysplasia			518	Acute myeloid leukemia M7
41	Acropigmentation of Dohi	466794	Acute infantile liver failure-cerebellar ataxia-peripheral sensory motor neuropathy syndrome	98831	Acute myeloid leukemia with 11q23 abnormalities
1133	Acrorenal defect-ectodermal dysplasia-diabetes syndrome				
971	Acrorenal syndrome	370088	Acute infantile liver failure-multisystemic involvement syndrome	98829	Acute myeloid leukemia with abnormal bone marrow eosinophils inv(16)(p13q22) or t(16;16)(p13;q22)
85203	ACRP syndrome				
36	ACS	98916	Acute inflammatory demyelinating polyradiculoneuropathy	319480	Acute myeloid leukemia with CEBPA somatic mutations
87	ACS1				
794	ACS3	98916	Acute inflammatory demyelinating polyneuropathy	402020	Acute myeloid leukemia with inv3(p21;q26.2) or t(3;3)(p21;q26.2)
710	ACS5				
98904	Actin myopathy	98916	Acute inflammatory polyneuropathy	98832	Acute myeloid leukemia with minimal differentiation
254395	Actinic lichen planus	79276	Acute intermittent porphyria	86845	Acute myeloid leukemia with multilineage dysplasia
254395	Actinic LP	79126	Acute interstitial pneumonia		
330061	Actinic prurigo	79126	Acute interstitial pneumonitis	402026	Acute myeloid leukemia with NPM1 somatic mutations
330064	Actinic reticuloid	73423	Acute intoxication by Blighia sapida		
457095	Actinomycosis	90062	Acute liver failure		
163696	Action myoclonus-renal failure syndrome				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
402014	Acute myeloid leukemia with t(6;9)(p23;q34)	43117	Acute tricyclic antidepressant poisoning	424991	Adenocarcinoma of gallbladder and extrahepatic biliary tract
370026	Acute myeloid leukemia with t(8;16)(p11;p13) translocation	91500	Acute tubulointerstitial nephritis and uveitis syndrome	424943	Adenocarcinoma of liver and IBT
102724	Acute myeloid leukemia with t(8;21)(q22;q22) translocation	98835	Acute undifferentiated leukemia	424943	Adenocarcinoma of liver and intrahepatic biliary tract
402017	Acute myeloid leukemia with t(9;11)(p22;q23)	284454	Acute zonal occult outer retinopathy	213504	Adenocarcinoma of ovary
		137754	ACY1D	363478	Adenocarcinoma of paratestis
		141	ACY2 deficiency	398053	Adenocarcinoma of penis
520	Acute myeloid leukemia with t(15;17)(q22;q12);(PML/RARalpha) and variants	99901	Acyl-CoA dehydrogenase 9 deficiency	104075	Adenocarcinoma of small bowel
517	Acute myelomonocytic leukemia	100008	ACys amyloidosis	104075	Adenocarcinoma of small intestine
86843	Acute myelosclerosis	99736	ACZ-responsive congenital myotonia	424016	Adenocarcinoma of the anal canal
263524	Acute necrotizing encephalopathy of childhood			95512	Adenohypophysitis
247546	Acute neonatal citrullinemia type 1	99736	ACZ-responsive myotonia	213828	Adenoid basal carcinoma of the cervix uteri
247546	Acute neonatal citrullinemia type I	93608	AD dRTA	213823	Adenoid cystic carcinoma of the cervix uteri
77260	Acute neuronopathic Gaucher disease	428	AD hypocalcemia	213741	Adenoid cystic carcinoma of the corpus uteri
		314889	AD pRTA		
		169189	AD-CNM		
163703	Acute non-herpetic encephalitis with severe refractory status epilepticus	1810	AD-HED	93292	Adenoma of pancreas
		2314	AD-HIES	26790	Adenomucinosi
		447753	AD-SPG9A	213792	Adenosarcoma of the cervix uteri
		447757	AD-SPG9B	213600	Adenosarcoma of the corpus uteri
35889	Acute opioid poisoning	277	ADA deficiency	45	Adenosine monophosphate deaminase deficiency
231457	Acute panautonomic GBS	435623	Adactyly of foot	28	Adenosylcobalamin deficiency
231457	Acute panautonomic Guillain-Barré syndrome	295118	Adactyly of foot, bilateral	91127	Adenovirus infection in immunocompromised patients
231457	Acute panautonomic neuropathy	295116	Adactyly of foot, unilateral		
231457	Acute pandysautonomia	295114	Adactyly of hand, bilateral	46	Adenylosuccinase deficiency
86843	Acute panmyelosis with myelofibrosis	973	Adactyly of hand, unilateral	46	Adenylosuccinate lyase deficiency
90064	Acute peripheral arterial occlusion	216796	Adair-Dighton syndrome	482601	Adenylosuccinate synthetase-like 1-related distal myopathy
43119	Acute poisoning by drugs with membrane-stabilizing effect	55881	Adamantinoma	137817	Adhesive arachnoiditis
520	Acute promyelocytic leukemia	55881	Adamantinoma of long bones	89937	ADHR
98918	Acute pure motor GBS	974	Adams-Oliver syndrome	454718	Adie syndrome
98918	Acute pure motor Guillain-Barré syndrome	97346	ADan amyloidosis	36397	Adiposalgia
231450	Acute pure sensory GBS	88619	ADANE	36397	Adipose tissue rheumatism
		314404	ADCA-DN syndrome	36397	Adiposis dolorosa
231450	Acute pure sensory Guillain-Barré syndrome	90348	ADCL	289290	ADK hypermethioninemia
231450	Acute pure sensory neuropathy	86814	ADCME	99027	ADLD
		85138	Addison disease	101046	ADLTE
231450	Acute pure sensory neuropathy	95409	Addisonian crisis	178464	ADMERF
454831	Acute radiation sickness	2953	Adducted thumb-clubfoot syndrome	98784	ADNFLE
454831	Acute radiation syndrome			329211	ADNIV
140896	Acute respiratory coronavirus infection	2952	Adducted thumbs-arthrogryposis syndrome, Christian type	404448	ADNP-related multiple congenital anomalies-intellectual disability-autism spectrum disorder
3099	Acute rheumatic fever	2953	Adducted thumbs-arthrogryposis syndrome, Dundar type		
90059	Acute sensorineural hearing loss by acute acoustic trauma or sudden deafness or surgery induced acoustic trauma	101046	ADEAF		
		83597	ADEM	306588	ADOS
231466	Acute sensory ataxic GBS	976	Adenine phosphoribosyltransferase deficiency	36355	ADP platelet receptor P2Y12 defect
231466	Acute sensory ataxic Guillain-Barré syndrome	213772	Adenocarcinoma of cervix uteri	2924	ADPCLD
		99976	Adenocarcinoma of esophagus	101046	ADPEAF
231466	Acute sensory ataxic neuropathy	424991	Adenocarcinoma of gallbladder and EBT	254892	adPEO
139417	Acute transverse myelitis			95409	Adrenal crisis

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
869	Adrenal insufficiency-achalasia-alacrima syndrome	99027	Adult-onset autosomal dominant leukodystrophy	3086	ADVIRC
1501	Adrenocortical carcinoma	284289	Adult-onset autosomal recessive cerebellar ataxia	682	Adynamia episodica hereditaria
231625	Adrenocortical carcinoma with pure aldosterone hypersecretion	255132	Adult-onset autosomal recessive sideroblastic anemia	1071	AEC syndrome
95409	Adrenocortical crisis	420492	Adult-onset cervical dystonia, DYT23 type	281139	AEI
99889	Adrenocorticotrophic hormone secretion syndrome	329336	Adult-onset chronic progressive external ophthalmoplegia with mitochondrial myopathy	163703	AERRPS
139399	Adrenomyeloneuropathy	247585	Adult-onset citrin deficiency	363549	AESD
977	Adrenomyodystrophy	247573	Adult-onset citrullinemia type 1	178345	AEXS
228169	ADSD	247573	Adult-onset citrullinemia type I	37	AEZ
46	ADSL deficiency	329336	Adult-onset CPEO with mitochondrial myopathy	220460	AFAP
482601	ADSSL1-related distal myopathy	411641	Adult-onset cystinosis	313772	AFG3L2-related spastic ataxia-neuropathy syndrome
70578	Adult acute respiratory distress syndrome	329478	Adult-onset distal myopathy due to VCP mutation	93562	AFib amyloidosis
70578	Adult ARDS	199351	Adult-onset dystonia-parkinsonism	243367	AFLP
93605	Adult Bartter syndrome	99000	Adult-onset foveomacular dystrophy	1827	AFND
157846	Adult basal ganglia disease	99000	Adult-onset foveomacular dystrophy with choroidal neovascularization	398147	AFP
874	Adult cardiac tumor	99000	Adult-onset foveomacular vitelliform dystrophy	139507	African iron overload
221	Adult dermatomyositis	79257	Adult-onset GM1 gangliosidosis	101334	African tick typhus
2666	Adult familial nephronophthisis-spastic quadriparesis syndrome	306431	Adult-onset immunodeficiency with anti-interferon-gamma autoantibodies	3385	African trypanosomiasis
309169	Adult GM2 gangliosidosis 0 variant	313808	Adult-onset leukoencephalopathy with axonal spheroids and pigmented glia	33110	Agammaglobulinemia, non-Bruton type
210159	Adult HCC	329314	Adult-onset multiple mitochondrial DNA deletion syndrome due to DGUOK deficiency	83617	Agammaglobulinemia-microcephaly-craniosynostosis-severe dermatitis syndrome
874	Adult heart tumor	329314	Adult-onset multiple mtDNA deletion syndrome due to DGUOK deficiency	388	Aganglionic megacolon
210159	Adult hepatocellular carcinoma	391490	Adult-onset myasthenia gravis	35704	AGAT deficiency
247676	Adult hypophosphatasia	171442	Adult-onset nemaline myopathy	85448	AGel amyloidosis
2688	Adult idiopathic neutropenia	276608	Adult-onset non-insulinoma persistent hyperinsulinemic hypoglycemia	180142	Agenesis and aplasia of uterine body
178487	Adult intestinal botulism	206572	Adult-onset overlap myositis	99114	Agenesis of the superior caval vein
178487	Adult intestinal colonization botulism	35689	Adult-onset PLS	99114	Agenesis of the superior vena cava
178487	Adult intestinal toxemia botulism	35689	Adult-onset primary lateral sclerosis	99114	Agenesis of the SVC
178487	Adult intestinal toxin-mediated botulism	209335	Adult-onset proximal spinal muscular atrophy, autosomal dominant	293173	AGEP
206448	Adult Krabbe disease	829	Adult-onset Still disease	873	Aggressive fibromatosis
79262	Adult NCL	247585	Adult-onset type 2 citrullinemia	86873	Aggressive NK-cell leukemia
79262	Adult neuronal ceroid lipofuscinosis	247585	Adult-onset type II citrullinemia	86873	Aggressive NK-cell lymphoma
247676	Adult phosphoethanolaminuria	247585	Adult-onset type II citrullinemia	98850	Aggressive systemic mastocytosis
206583	Adult polyglucosan body disease	99000	Adult-onset vitelliform macular dystrophy	989	Aglossia-adactylia syndrome
902	Adult progeria	99000	Adult-onset vitelliform macular dystrophy	990	Agnathia-holoprosencephaly-situs inversus syndrome
99874	Adult pulmonary Langerhans cell histiocytosis	99000	Adult-onset vitelliform macular dystrophy	824	Agnogenic myeloid metaplasia
98872	Adult pure red cell aplasia	99000	Adult-onset vitelliform macular dystrophy	100070	Agramatic variant of PPA
247676	Adult Rathburn disease	99000	Adult-onset vitelliform macular dystrophy	100070	Agramatic variant of primary progressive aphasia
773	Adult Refsum disease	99000	Adult-onset vitelliform macular dystrophy	442582	AH amyloidosis
978	ADULT syndrome	99000	Adult-onset vitelliform macular dystrophy	2131	AHC
86875	Adult T-cell leukemia/lymphoma	99000	Adult-onset vitelliform macular dystrophy	412069	AHDC1-related intellectual disability-obstructive sleep apnea-mild dysmorphism syndrome
391490	Adult-onset acquired myasthenia	99000	Adult-onset vitelliform macular dystrophy	59	AHDS
79280	Adult-onset Alpha-N-acetylgalactosaminidase deficiency	99000	Adult-onset vitelliform macular dystrophy	50812	Ahn-Lerman-Sagie syndrome
391490	Adult-onset autoimmune myasthenia gravis	99000	Adult-onset vitelliform macular dystrophy	79443	AHO-PHP syndrome Ia
99027	Adult-onset autosomal dominant demyelinating leukodystrophy	99000	Adult-onset vitelliform macular dystrophy	79445	AHO-PPHP syndrome
		99000	Adult-onset vitelliform macular dystrophy	2134	aHUS
		99000	Adult-onset vitelliform macular dystrophy	93581	aHUS with anti-factor H antibodies

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
93578	aHUS with B factor anomaly	261629	Alagille-Watson syndrome due to a NOTCH2 point mutation	79325	ALG8-CDG
93575	aHUS with C3 anomaly			79328	ALG9-CDG
357008	aHUS with DGKE deficiency	261600	Alagille-Watson syndrome due to monosomy 20p12	280071	ALG11-CDG
93579	aHUS with H factor anomaly			79324	ALG12-CDG
93580	aHUS with I factor anomaly	178333	Åland Islands eye disease	324422	ALG13-CDG
93576	aHUS with MCP/CD46 anomaly	2007	Alar cartilages hypoplasia-coloboma-telectanthis syndrome	99995	Algodystrophy
217023	aHUS with thrombomodulin anomaly	53	Albers-Schönberg osteopetrosis	300895	ALK+ ALCL
250977	AICA-ribosiduria	→897	Albinism-black lock-cell migration disorder of the neurocytes of the gut-sensorineural deafness syndrome	300895	ALK+ anaplastic large cell lymphoma
50	Aicardi syndrome			364043	ALK+ large B-cell lymphoma
51	Aicardi-Goutières syndrome			364043	ALK+ LBCL
101089	AID deficiency			300903	ALK- ALCL
98916	AIDP	998	Albinism-deafness syndrome	300903	ALK- anaplastic large cell lymphoma
90081	AIDS wasting syndrome	→457059	Albright hereditary osteodystrophy	300903	ALK-negative anaplastic large cell lymphoma
178333	AIED	1001	Albright hereditary osteodystrophy type 3	300895	ALK-positive anaplastic large cell lymphoma
363549	AIEF	1001	Albright hereditary osteodystrophy-like syndrome	364043	ALK-positive large B-cell lymphoma
86886	AILT			56	Alkaptonuria
103919	AIP	79443	Albright hereditary osteodystrophy-PHP syndrome Ia	59	Allan-Herndon-Dudley syndrome
280302	AIP type 1			1164	Allergic aspergillosis
280315	AIP type 2	79445	Albright hereditary osteodystrophy-PPHP syndrome	1164	Allergic bronchopulmonary aspergillosis
75564	AISA			869	Allgrove syndrome
33355	AK2 deficiency	98841	ALCL	69063	Alloimmune neonatal renal disease
38	AKE	60039	Alcock syndrome	93925	Alobar holoprosencephaly
→357225	Akesson syndrome	1915	Alcohol-related birth defects	1006	Alopecia antibody deficiency
79085	AKT2-related familial partial lipodystrophy	1915	Alcohol-related neurodevelopmental disorder	700	Alopecia totalis
79085	AKT2-related FPLD	36899	Alcohol-responsive dystonia	701	Alopecia universalis
85443	AL amyloidosis	43	ALD	2316	Alopecia-anosmia-deafness-hypogonadism syndrome
2232	Al Awadi-Farag-Teebi syndrome	324977	ALDD syndrome	1005	Alopecia-contractures-dwarfism-intellectual disability syndrome
2879	Al Awadi-Raas-Rothschild syndrome	35664	ALDH18A1-related De Barsy syndrome	202	Alopecia-deafness-hypogonadism syndrome
→3157	Al Frayh-Facharzt-Haque syndrome			2574	Alopecia-epilepsy-intellectual disability syndrome, Moynahan type
2725	Al Gazali-Al Talabani syndrome	99763	Aldosterone synthase deficiency	1008	Alopecia-epilepsy-pyorrhea-intellectual disability syndrome
2865	Al Gazali-Aziz-Salem syndrome	99764	Aldosterone synthase deficiency unrelated to CYP11B2	→3464	Alopecia-hypogonadism-extrapyramidal syndrome
2153	Al Gazali-Donnai-Muller syndrome			2850	Alopecia-intellectual disability syndrome
2725	Al Gazali-Lytle syndrome	99764	Aldosterone synthase deficiency unrelated to the aldosterone synthase gene	1014	Alopecia-intellectual disability-hypergonadotropic hypogonadism syndrome
2773	Al Gazali-Nair syndrome			157954	Alopecia-progressive neurological defect-endocrinopathy syndrome
→324737	Al-Gazali-Dattani syndrome	369929	Aldosterone-producing adenoma with seizures and neurological abnormalities	726	Alpers progressive sclerosing poliodystrophy
404454	Alacrimia-choreoathetosis-liver dysfunction syndrome			726	Alpers syndrome
100924	ALAD porphyria	369929	Aldosterone-secreting adenoma with seizures and neurological abnormalities	726	Alpers-Huttenlocher syndrome
52	Alagille syndrome			734	Alpha delta granule deficiency
261600	Alagille syndrome due to 20p12 microdeletion	85332	Aldred syndrome	734	Alpha dense granule deficiency
261619	Alagille syndrome due to a JAG1 point mutation	439224	ALECT2 amyloidosis	134	Alpha methylacetoacetic aciduria
261629	Alagille syndrome due to a NOTCH2 point mutation	158799	Aleukemic mast cell leukemia	721	Alpha storage pool deficiency
261600	Alagille syndrome due to del(20)(p12)	58	Alexander disease		
261600	Alagille syndrome due to monosomy 20p12	363717	Alexander disease type I		
52	Alagille-Watson syndrome	363722	Alexander disease type II		
261619	Alagille-Watson syndrome due to a JAG1 point mutation	261112	Alfi syndrome		
		79327	ALG1-CDG		
		79326	ALG2-CDG		
		79321	ALG3-CDG		
		79320	ALG6-CDG		

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98791	Alpha thalassemia-intellectual disability syndrome, deletion type	86818	Alport syndrome-intellectual disability-midface hypoplasia-elliptocytosis syndrome	100031	Amelogenesis imperfecta type 1
98791	Alpha thalassemia-mental retardation syndrome		3261	ALPS	100033
365	Alpha-1,4-glucosidase acid deficiency	436159	ALPS due to CTLA4 haploinsufficiency	100032	Amelogenesis imperfecta type 3
		268114	ALPS type 4	100034	Amelogenesis imperfecta type 4
308552	Alpha-1,4-glucosidase acid deficiency, infantile onset	436159	ALPS type 5	171836	Amelogenesis imperfecta-gingival hyperplasia syndrome
		268114	ALPS type IV		
420429	Alpha-1,4-glucosidase acid deficiency, late onset	436159	ALPS type V	1031	Amelogenesis imperfecta-nephrocalcinosis syndrome
		275517	ALPS with recurrent viral infections	1028	Amelonychohypohidrotic syndrome
60	Alpha-1-antitrypsin deficiency	803	ALS	83595	American mountain fever
79154	Alpha-aminoadipic aciduria	357043	ALS4	3386	American trypanosomiasis
399058	Alpha-B crystallin-related late-onset distal myopathy	86815	ALSG	2116	Aminoaciduria, Hartnup type
		313808	ALSP	141	Aminoacylase 2 deficiency
324	Alpha-galactosidase A deficiency	64	Alström syndrome	1908	Aminopterin embryopathy syndrome
100025	Alpha-HCD	99971	ALT	221120	Aminopterin syndrome-like sine aminopterin
100025	Alpha-heavy chain disease	2131	Alternating hemiplegia of childhood		
31	Alpha-ketoglutarate dehydrogenase deficiency	210122	Alveolar capillary dysplasia with misalignment of pulmonary veins	1908	Aminopterin/methotrexate embryofetopathy
349	Alpha-L-fucosidase deficiency	210122	Alveolar capillary dysplasia with misalignment of pulmonary vessels	→33364	Amish brittle hair syndrome
579	Alpha-L-iduronidase deficiency		Alveolar cleft lip and palate	171714	Amish infantile epilepsy syndrome
61	Alpha-mannosidosis	199306	Alveolar rhabdomyosarcoma	99742	Amish lethal microcephaly
309288	Alpha-mannosidosis, adult form	284	Alveolar soft-part sarcoma	98902	Amish nemaline myopathy
309282	Alpha-mannosidosis, infantile form	99756	Alveolar soft-tissue sarcoma	518	AMKL
		163699	Alveolar synechia-ankyloblepharon-ectodermal dysplasia syndrome	102379	AML and myelodysplastic syndromes related to alkylating agent
134	Alpha-methyl-acetoacetyl-CoA thiolase deficiency	163699	Alves-dos Santos-Castelo syndrome	164726	AML and myelodysplastic syndromes related to radiation
79095	Alpha-methyl-acyl-CoA racemase deficiency	→1071	ALX1-related frontonasal dysplasia	102381	AML and myelodysplastic syndromes related to topoisomerase type 2 inhibitor
3137	Alpha-N-acetylgalactosaminidase deficiency	3354	ALX3-related frontonasal dysplasia		
		306542	ALX4-related FNDAG		
79279	Alpha-N-acetylgalactosaminidase deficiency type 1	391474	Alymphoid cystic thymic dysgenesis	98832	AML M0
79280	Alpha-N-acetylgalactosaminidase deficiency type 2	228390	ALys amyloidosis	98833	AML M1
		169095	AMACR deficiency	98834	AML M2
79281	Alpha-N-acetylgalactosaminidase deficiency type 3	93561	AMAN	520	AML M3
62	Alpha-sarcoglycanopathy	98918	Amaurosis congenita of Leber	517	AML M4
846	Alpha-thalassemia	65	Amaurosis-hypertrichosis syndrome	514	AML M5
163596	Alpha-thalassemia hydrops fetalis	1021	Ambiguous genitalia-disordered steroidogenesis Antley-Bixler-like syndrome	318	AML M6
93616	Alpha-thalassemia intermedia	→95699	Ambras syndrome	518	AML M7
163596	Alpha-thalassemia major			98831	AML with 11q23 abnormalities
98791	Alpha-thalassemia-intellectual disability syndrome linked to chromosome 16	1023	Amelia of lower limb	98829	AML with abnormal bone marrow eosinophils inv(16)(p13q22) or t(16;16)(p13;q22)
		294969	Amelia of lower limb, bilateral		
231401	Alpha-thalassemia-myelodysplastic syndrome	295059	Amelia of lower limb, unilateral	319480	AML with CEBPA somatic mutations
		295057	Amelia of upper limb	402020	AML with inv3(p21;q26.2) or t(3;3)(p21;q26.2)
847	Alpha-thalassemia-X-linked intellectual disability syndrome	294967	Amelia of upper limb, bilateral		
63	Alport deafness-nephropathy	295055	Amelia of upper limb, unilateral	86845	AML with multilineage dysplasia
63	Alport syndrome	295053	Ameloblastic carcinoma	402026	AML with NPM1 somatic mutations
		314422	Ameloblastoma	402014	AML with t(6;9)(p23;q34)
→182050	Alport syndrome with leukocyte inclusions and macrothrombocytopenia	314419	Ameloblastoma of jaw	370026	AML with t(8;16)(p11;p13) translocation
		314419	Amelocerebrohypohidrotic syndrome		
→182050	Alport syndrome with macrothrombocytopenia	1946	Amelogenesis imperfecta	102724	AML with t(8;21)(q22;q22) translocation
		88661			

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402017	AML with t(9;11)(p22;q23)	37553	Andersen syndrome		
520	AML with t(15;17)(q22;q12);(PML/RARalpha) and variants	37553	Andersen-Tawil syndrome	69088	Anhidrotic ectodermal dysplasia-immunodeficiency-osteopetrosis-lymphedema syndrome
		71	Anderson disease		
		324	Anderson-Fabry disease	→1071	Anhidrotic ectodermic dysplasia-cleft lip/palate syndrome
86818	AMME complex	99916	Androblastoma		
86818	AMME syndrome	329813	Androgenetic/biparental mosaicism	1069	Aniridia-absent patella syndrome
517	AMMoL	157954	ANE syndrome	1065	Aniridia-cerebellar ataxia-intellectual disability syndrome
251663	aMOA	263524	ANEC		
67	Amoebiasis due to Entamoeba histolytica	1044	Anemia due to adenosine triphosphatase deficiency	1068	Aniridia-intellectual disability syndrome
68	Amoebiasis due to free-living amoebae	1054	Aneurysm of sinus of Valsalva	1067	Aniridia-ptosis-intellectual disability-familial obesity syndrome
45	AMP deaminase deficiency	95484	Aneurysm or dilatation of ascending aorta	1064	Aniridia-renal agenesis-psychomotor retardation syndrome
1035	Ampola syndrome	284984	Aneurysm-osteoarthritis syndrome		
66529	Ampulla cardiomyopathy	480553	Aneurysmal bone cyst	1070	Anisakiasis
300557	Ampullary carcinoma	353344	Aneurysmal telangiectasia	86873	ANKCL
300557	Ampulloma			1072	Ankyloblepharon filiforme adnatum
98917	AMSAN	63442	Angel-shaped phalango-epiphyseal dysplasia	1072	Ankyloblepharon filiforme adnatum-cleft palate syndrome
366	Amylo-1,6-glucosidase deficiency	72	Angelman syndrome		
49804	Amyloid lichen			1074	Ankyloblepharon filiforme-imperforate anus syndrome
319635	Amyloidosis cutis dyschromia	411511	Angelman syndrome due to a point mutation		
319635	Amyloidosis cutis dyschromica	411515	Angelman syndrome due to imprinting defect in 15q11-q13	1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome
85450	Amyloidosis, Ostertag type			2206	Ankylosing vertebral hyperostosis with tylosis
367	Amylopectinosis	98794	Angelman syndrome due to maternal 15q11q13 deletion		
803	Amyotrophic lateral sclerosis			1077	Ankylosis of teeth
357043	Amyotrophic lateral sclerosis type 4	98794	Angelman syndrome due to maternal monosomy 15q11q13	78	Ankylostomiasis
94091	Amyotrophic lateral sclerosis, hemiplegic type			254411	Annular atrophic lichen planus
90020	Amyotrophic lateral sclerosis-parkinsonism-dementia complex	98795	Angelman syndrome due to paternal uniparental disomy of chromosome 15	254411	Annular atrophic LP
90020	Amyotrophic lateral sclerosis-parkinsonism-dementia of Guam syndrome	251671	Angiocentric glioma	281139	Annular epidermolytic ichthyosis
		86879	Angiocentric T-cell lymphoma	254424	Annular lichen planus
		79093	Angiodysgenetic necrotizing myelopathy	254424	Annular LP
2615	Amyotrophy-fat tissue anomaly syndrome	98839	Angioendotheliomatosis proliferans systemisata	675	Annular pancreas
228113	Anal fistula			229	Annuloaortic ectasia
31150	Analphalipoproteinemia	160	Angiofollicular ganglionic hyperplasia	457205	ANOAC
761	Anaphylactoid purpura			99797	Anodontia
251589	Anaplastic astrocytoma	160	Angiofollicular lymph hyperplasia	101932	Anomaly of the mitral subvalvular apparatus
251646	Anaplastic ependymoma	86886	Angioimmunoblastic T-cell lymphoma	99055	Anomaly of the tricuspid valve chordae
251957	Anaplastic ganglioglioma			94150	Anonychia congenita totalis
98841	Anaplastic large cell lymphoma	324	Angiokeratoma corporis diffusum	69125	Anonychia with flexural pigmentation
251663	Anaplastic oligoastrocytoma	95429	Angioma serpiginosum		
251630	Anaplastic oligodendroglioma	2346	Angioosteohypertrophic syndrome	1094	Anonychia-microcephaly syndrome
142	Anaplastic thyroid carcinoma	75508	Angioosteohypotrophic syndrome		
251855	Anaplastic/large cell medulloblastoma	263413	Angiosarcoma	90390	Anonychia-onychodystrophy syndrome
		74	Angiostrongyliasis		
93347	Anauxetic dysplasia	98839	Angiotropic large cell lymphoma	1487	Anonychia-onychodystrophy with hypoplasia or absence of distal phalanges syndrome
79262	ANCL	370039	Angora hair nevus		
78	Ancylostomiasis	76	Anguilluliasis	1104	Anopthalmia plus syndrome
1496	Andermann syndrome	76	Anguillulosis		
37553	Andersen cardiodysrhythmic periodic paralysis	238468	Anhidrotic ectodermal dysplasia	→2470	Anopthalmia-heart and pulmonary anomalies-intellectual disability syndrome
		98813	Anhidrotic ectodermal dysplasia with immunodeficiency	→3157	Anopthalmia-hypothalamo-pituitary insufficiency syndrome
367	Andersen disease				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
1101	Anophthalmia-megalocornea-cardiopathy-skeletal anomalies syndrome	→95699	Antley-Bixler syndrome type 2	1113	Aphalangy-syndactyly-microcephaly syndrome
		→95699	Antley-Bixler syndrome with genital anomaly and disorder of steroidogenesis	49	Aphallia
2470	Anophthalmia-pulmonary hypoplasia syndrome	→95699	Antley-Bixler syndrome, POR-related	324540	Aphonia-deafness-retinal dystrophy-bifid halluces-intellectual disability syndrome
1106	Anophthalmia-syndactyly syndrome			1190	AO1
77298	Anophthalmia/microphthalmia-esophageal atresia syndrome	56304	AO2	66529	Apical ballooning syndrome
1882	ANOTHER syndrome	56305	AO3		
93976	Anotia	1168	AOA1	1114	Aplasia cutis congenita
2987	Antecubital pterygium syndrome	64753	AOA2	3339	Aplasia cutis congenita-epibulbar dermoids syndrome
93604	Antenatal Bartter syndrome	459033	AOA4	1116	Aplasia cutis congenita-intestinal lymphangiectasia syndrome
294	Antenatal CMV infection	99000	AOFMD	370046	Aplasia cutis congenita-nevus sebaceus syndrome
294	Antenatal cytomegalovirus infection	1190	AOI		
70596	Antenatal EBV infection	56304	AOII	86815	Aplasia of lacrimal and salivary glands
292	Antenatal enterovirus infection	56305	AOIII	3329	Aplasia of tibia with split-hand/split-foot deformity
70596	Antenatal Epstein-Barr virus infection	1457	Aorta coarctation	2879	Aplasia/hypoplasia of limbs and pelvis
		60030	Aortic aneurysm syndrome due to TGF-beta receptors anomalies	520	APML
293	Antenatal herpes simplex virus infection	1110	Aortic arch anomaly-facial dysmorphism-intellectual disability syndrome	70590	Apnea of infancy
178148	Antenatal multimimicore disease with arthrogryposis multiplex congenita			2299	Aortic arch interruption
		99079	Aortic arch syndrome	425	ApoA-I deficiency
291	Antenatal varicella virus infection	→91387	Aortic dilatation-joint hypermobility-arterial tortuosity syndrome	294986	Apodia
51890	Anterior cutaneous nerve entrapment syndrome	95448	Aortic valve atresia	295107	Apodia, bilateral
1931	Anterior encephalocele	101043	Aortic valve dysplasia	295105	Apodia, unilateral
98961	Anterior limiting membrane dystrophy type 1	99071	Aorto-left ventricular tunnel	93560	Apolipoprotein A-I amyloidosis
		99070	Aorto-right ventricular tunnel	425	Apolipoprotein A-I deficiency
98961	Anterior limiting membrane dystrophy type I	3400	Aorto-ventricular tunnel	238269	Apolipoprotein A-II amyloidosis
98960	Anterior limiting membrane dystrophy type II	99086	Aortopulmonary coronary arterial course	439232	Apolipoprotein A-IV amyloidosis
95512	Anterior pituitary hypophysitis	974	AOS	320	Apparent mineralocorticoid excess
435372	Anterior urethral valve	829	AOSD	391723	Appendiceal mucinous adenocarcinoma
90079	Anthracycline extravasation	280763	AP4 deficiency syndrome	100079	Appendiceal neuroendocrine tumor
36412	Anti-C1q vasculitis	369929	APA with seizures and neurological abnormalities	1201	Apple peel syndrome
375	Anti-GBM syndrome	747	aPAP	1126	Aprosencephaly cerebellar dysgenesis
375	Anti-glomerular basement membrane disease	206583	APBD	976	APRT deficiency
2194	Anti-HLA hyperimmunization	247806	APC-related AFAP	3453	APS1
206569	Anti-HMG-CoA myopathy	247806	APC-related attenuated familial adenomatous polyposis	3143	APS2
81	Anti-Jo1 syndrome	247806	APC-related attenuated familial polyposis coli	227982	APS3
639	Anti-MAG neuropathy			227990	APS4
454710	Anti-p200 pemphigoid	247806	APC-related attenuated FAP	3453	APS type 1
206569	Anti-SRP myopathy	397596	APDS	3143	APS type 2
97564	Antineutrophil cytoplasmic antibody-negative pauci-immune glomerulonephritis	3453	APECED syndrome	227982	APS type 3
		87	Apert syndrome	227990	APS type 4
2821	Antinolo-Nieto-Borrego syndrome	162521	Apertura pyriformis with holoprosencephaly	101206	APV/ADA, Fallot type
3006	Antiquitin deficiency	1112	Aphalangy-hemivertebrae-urogenital-intestinal dysgenesis syndrome	99048	APV/PDA, non-Fallot type
81	Antisynthetase syndrome			402041	AR dRTA
83	Antley-Bixler syndrome				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
→402041	AR dRTA with deafness	247525	Argininosuccinic acid synthase deficiency	1150	Arthrogryposis multiplex congenita-whistling face syndrome
→402041	AR dRTA with hearing loss				
→402041	AR dRTA without deafness				
→402041	AR dRTA without hearing loss				
93607	AR pRTA	23	Argininosuccinic aciduria	1154	Arthrogryposis-oculomotor limitation-electroretinal anomalies syndrome
90119	AR-CMT2 with acrodystrophy	60014	Argyria		
90118	AR-CMT2, Ouvrier type	289176	ARHR		
98856	AR-CMT2B1	79235	Arias syndrome		
101101	AR-CMT2B2	2318	Arima syndrome		
101102	AR-CMT2C	950	Arkless-Graham syndrome		
443950	AR-CMT2T	85276	Armfield syndrome		
169186	AR-CNM	1915	ARND		
248	AR-HED	167635	Arndt-Gottron disease		
88616	AR-NSID	268882	Arnold-Chiari malformation type 1		
731	AR-PKD	1136	Arnold-Chiari malformation type 2		
447760	AR-SPG9B	268882	Arnold-Chiari malformation type I		
1129	Arachnodactyly-abnormal ossification-intellectual disability syndrome	1136	Arnold-Chiari malformation type II		
		91	Aromatase deficiency		
		178345	Aromatase excess syndrome		
1130	Arachnodactyly-intellectual disability-dysmorphism syndrome	35708	Aromatic L-amino acid decarboxylase deficiency	65720	Arthrogryposis-severe scoliosis syndrome
2356	Arachnoid cyst	254886	arPEO		
137817	Arachnoiditis	99916	Arrhenoblastoma	2848	Arthropathy-camptodactyly syndrome
324442	ARAN-NM	1134	Arrhinia		
1915	ARBD	1135	Arrhinia-choanal atresia-microphthalmia syndrome	309271	Arylsulfatase A deficiency, adult form
2697	ARC syndrome				
88644	ARCA1	260305	ARSA	309263	Arylsulfatase A deficiency, juvenile form
139485	ARCA2	98	ARSACS		
90349	ARCL1	314603	ARSAL	309256	Arylsulfatase A deficiency, late infantile form
357074	ARCL2, classic type	583	ARSB deficiency		
357074	ARCL2, Debré type	357107	Arterial cervical rib syndrome	583	Arylsulfatase B deficiency
357064	ARCL2, progeroid type	357107	Arterial costoclavicular syndrome		
357058	ARCL2A	1682	Arterial dissection-lentiginosis syndrome	276212	Arylsulfatase B deficiency, rapidly progressing
357064	ARCL2B				
324442	ARCMT2-NM	357107	Arterial hyperabduction syndrome	276223	Arylsulfatase B deficiency, slowly progressing
101097	ARCMT2K	357107	Arterial scalenus anticus syndrome		
466775	ARCMT2X	357107	Arterial thoracic outlet compression syndrome	81	AS syndrome
1133	AREDYLD syndrome				
101096	Aregenerative anemia	3342	Arterial tortuosity syndrome	23	ASA deficiency
→702	Arena syndrome				
75377	Areolar atrophy of the macula	357107	Arterial TOS	231466	ASAN
319223	Argentine hemorrhagic fever				
319223	Argentinian hemorrhagic fever	52	Arteriohepatic dysplasia	583	ASB deficiency
90	Arginase deficiency				
90	Argininemia	261619	Arteriohepatic dysplasia due to a JAG1 point mutation	2302	Asbestos intoxication
23	Argininosuccinase deficiency				
247525	Argininosuccinate synthase deficiency	261629	Arteriohepatic dysplasia due to a NOTCH2 point mutation	2302	Asbestosis
247525	Argininosuccinate synthetase deficiency				
23	Argininosuccinatelyase deficiency	261600	Arteriohepatic dysplasia due to monosomy 20p12	1253	Ascher syndrome
23	Argininosuccinic acid lyase deficiency	29207	Arthritis urethritica		
		955	Arthrodentoosteodysplasia		
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	1155	Arthrogryposis due to muscular dystrophy	447997	ASCT1 deficiency
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	1478	ASD
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	352490	ASD due to AUTS2 deficiency
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	99104	ASD, coronary sinus type
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	99106	ASD, ostium primum type
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	99103	ASD, ostium secundum type
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	99105	ASD, sinus venosus type
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	54251	Aseptic abscesses syndrome
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	97337	Aseptic necrosis of patella
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	3314	Aseptic necrosis of phalangeal epiphyses
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	2380	Aseptic necrosis of the capital femoral epiphysis
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	97336	Aseptic necrosis of the capital humerus
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	97332	Aseptic necrosis of the lunate bone
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome	2054	Aseptic necrosis of the tarsal bone
994	Arthrogryposis multiplex congenita-pulmonary hypoplasia syndrome				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
97335	Aseptic necrosis of the tibial tubercle	56304	Atelosteogenesis type 2	1342	Atrioidigital dysplasia type 3
		56305	Atelosteogenesis type 3	168796	Atrioidigital dysplasia, Slovenian type
57194	Aseptic osteitis	1190	Atelosteogenesis type I		
54251	Aseptic systemic abscesses	56304	Atelosteogenesis type II	1352	Atrioventricular defect-blepharophimosis-radial and anal defect syndrome
137686	Asherman syndrome	56305	Atelosteogenesis type III		
276198	Asidan	69739	Athabaskan brainstem dysgenesis syndrome	86813	Atrophia areata
23	ASL deficiency			649	Atrophia bulborum hereditaria
391376	Asparagine synthetase deficiency	69739	Athabaskan brainstem dysgenesis syndrome	254449	Atrophic lichen planus
141	Aspartoacylase deficiency			254449	Atrophic LP
93	Aspartylglucosaminidase deficiency	1192	Atherosclerosis-deafness-diabetes-epilepsy-nephropathy syndrome	79100	Atrophoderma vermiculata
93	Aspartylglucosaminuria			99966	ATRT
63442	ASPED	95713	Athyreosis	71289	ATRUS syndrome
1163	Aspergillosis	1226	Athyroidal hypothyroidism-spiky hair-cleft palate syndrome	3342	ATS
474	Asphyxiating thoracic dystrophy of the newborn	250977	ATIC deficiency	86818	ATS-MR
163699	ASPS	1193	Atkin-Flaitz syndrome	352723	Attenuated Chédiak-Higashi syndrome
247525	ASS deficiency	99666	Atlantoaxial subluxation		
221120	ASSA	251347	ATLD	220460	Attenuated familial adenomatous polyposis
85175	Astley-Kendall dysplasia	86875	ATLL	220460	Attenuated familial polyposis coli
251679	Astroblastoma	139423	ATM/TM	220460	Attenuated FAP
647	AT V1	231401	ATMDS	85451	ATTR cardiomyopathy
96	Ataxia with isolated vitamin E deficiency	163934	Atopic keratoconjunctivitis	85451	ATTRV122I amyloidosis
		357107	ATOS	85451	ATTRV122I-related amyloidosis
3008	Ataxia with lactic acidosis type 2	139557	ATP7A-related distal motor neuropathy	85447	ATTRV30M amyloidosis
3008	Ataxia with lactic acidosis type II			85447	ATTRV30M-related amyloidosis
94147	Ataxia with pigmentary retinopathy	31150	ATP-binding cassette transporter A1 deficiency	330001	ATTRwt amyloidosis
96	Ataxia with vitamin E deficiency			330001	ATTRwt-related amyloidosis
1188	Ataxia-deafness-intellectual disability syndrome	98791	ATR syndrome linked to chromosome 16	199627	Atypical autism
		98791	ATR syndrome, deletion type	352723	Atypical Chédiak-Higashi syndrome
137639	Ataxia-delayed dentition-hypomyelination syndrome	98791	ATR-16 syndrome	251902	Atypical choroid plexus papilloma
		847	ATR-X syndrome	98824	Atypical chronic myeloid leukemia
1227	Ataxia-diabetes-goiter-gonadal insufficiency syndrome	30391	Atresia of bile ducts	1456	Atypical coarctation of aorta
		1201	Atresia of small intestine	251902	Atypical CPP
1180	Ataxia-hypogonadism-choroidal dystrophy syndrome	105	Atresia of urethra	314466	Atypical Demons-Meigs syndrome
		1344	Atrial cardiomyopathy with heart block	314721	Atypical dentin dysplasia due to SMOC2 deficiency
370022	Ataxia-intellectual disability-oculomotor apraxia-cerebellar cysts syndrome	99107	Atrial septal aneurysm	398147	Atypical facial pain
1168	Ataxia-oculomotor apraxia type 1	1478	Atrial septal defect	309252	Atypical Gaucher disease due to saposin C deficiency
64753	Ataxia-oculomotor apraxia type 2	99104	Atrial septal defect, coronary sinus type	289863	Atypical glycine encephalopathy
459033	Ataxia-oculomotor apraxia type 4			98961	Atypical granular corneal dystrophy
2585	Ataxia-pancytopenia syndrome	99106	Atrial septal defect, ostium primum type	238523	Atypical HCS
1184	Ataxia-photosensitivity-short stature syndrome	99103	Atrial septal defect, ostium secundum type	2134	Atypical hemolytic-uremic syndrome
1178	Ataxia-tapetoretinal degeneration syndrome	99105	Atrial septal defect, sinus venosus type	93581	Atypical hemolytic-uremic syndrome with anti-factor H antibodies
100	Ataxia-telangiectasia	1479	Atrial septal defect-atrioventricular conduction defects syndrome	93578	Atypical hemolytic-uremic syndrome with B factor anomaly
370109	Ataxia-telangiectasia variant	1344	Atrial standstill	93575	Atypical hemolytic-uremic syndrome with C3 anomaly
647	Ataxia-telangiectasia, variant 1	844	Atrial tachyarrhythmia with short PR interval		
251347	Ataxia-telangiectasia-like disorder	86819	Atrichia with papular lesions	357008	Atypical hemolytic-uremic syndrome with DGKE deficiency
1183	Ataxo-opsomyoclonus syndrome	392	Atrioidigital dysplasia type 1		
2953	ATCS	1350	Atrioidigital dysplasia type 2		
3469	Atelencephaly				
1190	Atelosteogenesis type 1				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
93579	Atypical hemolytic-uremic syndrome with H factor anomaly	1488	Aural atresia-multiple congenital anomalies-intellectual disability syndrome	436159	Autoimmune lymphoproliferative syndrome type 5
93580	Atypical hemolytic-uremic syndrome with I factor anomaly		→794	Auralcephalosyndactyly	268114
93576	Atypical hemolytic-uremic syndrome with MCP/CD46 anomaly	77300	Auricular abnormalities-cleft lip with or without cleft palate-ocular abnormalities syndrome	436159	Autoimmune lymphoproliferative syndrome type V
217023	Atypical hemolytic-uremic syndrome with thrombomodulin anomaly	137888	Auriculocondylar syndrome	275517	Autoimmune lymphoproliferative syndrome with recurrent viral infections
		114	Auriculoosteodysplasia		
2134	Atypical HUS	→794	Aurocephalosyndactyly	589	Autoimmune myasthenia gravis
93581	Atypical HUS with anti-factor H antibodies	1995	Ausems-Wittebol Post-Hennekam syndrome	206569	Autoimmune necrotizing myositis
93578	Atypical HUS with B factor anomaly	352490	Autism spectrum disorder due to AUTS2 deficiency	103919	Autoimmune pancreatitis
93575	Atypical HUS with C3 anomaly			280302	Autoimmune pancreatitis type 1
357008	Atypical HUS with DGKE deficiency	370943	Autism spectrum disorder-epilepsy-arthrogryposis syndrome	280315	Autoimmune pancreatitis type 2
93579	Atypical HUS with H factor anomaly			747	Autoimmune PAP
93580	Atypical HUS with I factor anomaly	308410	Autism-epilepsy syndrome due to branched chain ketoacid dehydrogenase kinase deficiency	3453	Autoimmune polyendocrine syndrome type 1
93576	Atypical HUS with MCP/CD46 anomaly			3143	Autoimmune polyendocrine syndrome type 2
217023	Atypical HUS with thrombomodulin anomaly	137911	Autism-facial port-wine stain syndrome	227982	Autoimmune polyendocrine syndrome type 3
238523	Atypical hypotonia-cystinuria syndrome	324636	Autoerythrocyte sensitization syndrome	227990	Autoimmune polyendocrine syndrome type 4
391411	Atypical juvenile parkinsonism	85138	Autoimmune Addison's disease	3453	Autoimmune polyendocrinopathy type 1
86797	Atypical lichen myxedematosus	85138	Autoimmune adrenalitis		
99971	Atypical lipoma	420789	Autoimmune encephalopathy with parasomnia and obstructive sleep apnea	3143	Autoimmune polyendocrinopathy type 2
99971	Atypical lipomatous tumor			227982	Autoimmune polyendocrinopathy type 3
314466	Atypical Meigs syndrome	391487	Autoimmune enteropathy and endocrinopathy-susceptibility to chronic infections syndrome	227990	Autoimmune polyendocrinopathy type 4
2578	Atypical MRKH syndrome			37042	Autoimmune enteropathy type 1
289863	Atypical NKA	103916	Autoimmune enteropathy type 2	3453	Autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy syndrome
289863	Atypical non-ketotic hyperglycinemia	103917	Autoimmune enteropathy type 3		
261501	Atypical Norrie disease due to del(X)(p11.3)	1959	Autoimmune hemolytic anemia and autoimmune thrombocytopenia	3453	Autoimmune polyglandular syndrome type 1
261501	Atypical Norrie disease due to monosomy Xp11.3	90033	Autoimmune hemolytic anemia, warm type	3143	Autoimmune polyglandular syndrome type 2
216873	Atypical pantothenate kinase-associated neurodegeneration	444463	Autoimmune hemolytic anemia-autoimmune thrombocytopenia-primary immunodeficiency syndrome	227982	Autoimmune polyglandular syndrome type 3
251902	Atypical papilloma of choroid plexus			227990	Autoimmune polyglandular syndrome type 4
79474	Atypical progeroid syndrome	2137	Autoimmune hepatitis	747	Autoimmune pulmonary alveolar proteinosis
99750	Atypical progressive supranuclear palsy syndrome	36913	Autoimmune hypoparathyroidism		
99750	Atypical PSP syndrome	3453	Autoimmune hypoparathyroidism-chronic candidiasis-Addison disease syndrome	93585	Autoimmune thrombotic thrombocytopenic purpura
3095	Atypical Rett syndrome			3143	Autoimmune thyroid disease and/or type 1 diabetes-Addison disease syndrome
3095	Atypical RTT	444092	Autoimmune interstitial lung disease-arthritis syndrome	324977	Autoinflammation-lipodystrophy-dermatosis syndrome
99966	Atypical teratoid rhabdoid tumor				
90393	Atypical tuberous myxedema of Jadassohn-Dosseker	3261	Autoimmune lymphoproliferative syndrome	324530	Autoinflammation-PLCG2-associated antibody deficiency-immune dysregulation
79474	Atypical Werner syndrome	436159	Autoimmune lymphoproliferative syndrome due to CTLA4 haploinsufficiency		
16	Atypical X-linked achromatopsia	268114	Autoimmune lymphoproliferative syndrome type 4		
453504	Au-Kline syndrome				
166415	Audiogenic seizures				
1074	Aughton-Hufnagle syndrome				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
210115	Autoinflammatory disease due to interleukin-1 receptor antagonist deficiency	99939	Autosomal dominant Charcot-Marie-Tooth disease type 2E	75381	Autosomal dominant cystoid macular edema
329173	Autoinflammatory syndrome with pyogenic bacterial infection and amylopectinosis	99940	Autosomal dominant Charcot-Marie-Tooth disease type 2F	79499	Autosomal dominant deafness-onychodystrophy syndrome
		99941	Autosomal dominant Charcot-Marie-Tooth disease type 2G	2337	Autosomal dominant diffuse palmoplantar keratoderma, Norrbotten type
33110	Autosomal agammaglobulinemia	99942	Autosomal dominant Charcot-Marie-Tooth disease type 2I	476093	Autosomal dominant distal axonal motor neuropathy-myofibrillar myopathy syndrome
88918	Autosomal dominant Alport syndrome	99943	Autosomal dominant Charcot-Marie-Tooth disease type 2J		139518
1810	Autosomal dominant anhidrotic ectodermal dysplasia	99944	Autosomal dominant Charcot-Marie-Tooth disease type 2K	93608	Autosomal dominant distal renal tubular acidosis
314399	Autosomal dominant aplasia and myelodysplasia	99945	Autosomal dominant Charcot-Marie-Tooth disease type 2L	98808	Autosomal dominant dopa-responsive dystonia
314399	Autosomal dominant aplastic anemia and myelodysplasia	228179	Autosomal dominant Charcot-Marie-Tooth disease type 2M	→231568	Autosomal dominant dystrophic epidermolysis bullosa, Cockayne-Touraine type
1216	Autosomal dominant benign distal spinal muscular atrophy	228174	Autosomal dominant Charcot-Marie-Tooth disease type 2N		231568
314652	Autosomal dominant beta2-microglobulinic amyloidosis	284232	Autosomal dominant Charcot-Marie-Tooth disease type 2O	→231568	Autosomal dominant dystrophic epidermolysis bullosa, Pasini type
93304	Autosomal dominant brachyolmia	329258	Autosomal dominant Charcot-Marie-Tooth disease type 2Q		300576
169189	Autosomal dominant centronuclear myopathy	397735	Autosomal dominant Charcot-Marie-Tooth disease type 2U	98853	Autosomal dominant Emery-Dreifuss muscular dystrophy
314404	Autosomal dominant cerebellar ataxia-deafness-narcolepsy syndrome	447964	Autosomal dominant Charcot-Marie-Tooth disease type 2V	101046	Autosomal dominant epilepsy with auditory features
		466768	Autosomal dominant Charcot-Marie-Tooth disease type 2Z	73229	Autosomal dominant familial hematuria-retinal arteriolar tortuosity-contractions syndrome
397735	Autosomal dominant Charcot-Marie-Tooth disease type 2 due to MARS mutation	98975	Autosomal dominant CHED	100988	Autosomal dominant familial spastic paraplegia type 3
466768	Autosomal dominant Charcot-Marie-Tooth disease type 2 due to MORC2 mutation	363447	Autosomal dominant childhood-onset proximal spinal muscular atrophy	329466	Autosomal dominant focal dystonia, DYT25
447964	Autosomal dominant Charcot-Marie-Tooth disease type 2 due to NAGLU mutation	363454	Autosomal dominant childhood-onset proximal spinal muscular atrophy with contractures	402003	Autosomal dominant focal non-epidermolytic palmoplantar keratoderma with plantar blistering
435819	Autosomal dominant Charcot-Marie-Tooth disease type 2 due to TFG mutation	209341	Autosomal dominant childhood-onset proximal spinal muscular atrophy without contractures	2024	Autosomal dominant gingival fibromatosis
435387	Autosomal dominant Charcot-Marie-Tooth disease type 2 due to VCP mutation	79344	Autosomal dominant chondrodysplasia punctata	2024	Autosomal dominant gingival hyperplasia
401964	Autosomal dominant Charcot-Marie-Tooth disease type 2 with giant axons	→2526	Autosomal dominant chorioretinopathy-microcephaly syndrome	139491	Autosomal dominant hereditary hemochromatosis
99946	Autosomal dominant Charcot-Marie-Tooth disease type 2A1	1455	Autosomal dominant coarctation of aorta	401964	Autosomal dominant hereditary motor and sensory neuropathy type 2 with giant axons
99947	Autosomal dominant Charcot-Marie-Tooth disease type 2A2	447753	Autosomal dominant complex spastic paraplegia type 9A	2314	Autosomal dominant HIES
99936	Autosomal dominant Charcot-Marie-Tooth disease type 2B	447757	Autosomal dominant complex spastic paraplegia type 9B	2314	Autosomal dominant hyper-IgE syndrome
99937	Autosomal dominant Charcot-Marie-Tooth disease type 2C	1216	Autosomal dominant congenital benign spinal muscular atrophy	2314	Autosomal dominant hyperimmunoglobulin E syndrome
99938	Autosomal dominant Charcot-Marie-Tooth disease type 2D	98975	Autosomal dominant congenital hereditary endothelial dystrophy	2314	Autosomal dominant hyperimmunoglobulin E syndrome
		86814	Autosomal dominant cortical myoclonus and epilepsy		
		90348	Autosomal dominant cutis laxa		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
276580	Autosomal dominant hyperinsulinemic hypoglycemia due to Kir6.2 deficiency	411602	Autosomal dominant late-onset Parkinson disease	319589	Autosomal dominant MSMD due to partial IFN $\gamma$ R2 deficiency
276575	Autosomal dominant hyperinsulinemic hypoglycemia due to SUR1 deficiency	67042	Autosomal dominant late-onset retinal degeneration	319581	Autosomal dominant MSMD due to partial interferon gamma receptor 1 deficiency
		101046	Autosomal dominant lateral temporal lobe epilepsy	319589	Autosomal dominant MSMD due to partial interferon gamma receptor 2 deficiency
276580	Autosomal dominant hyperinsulinism due to Kir6.2 deficiency	313808	Autosomal dominant leukoencephalopathy with neuroaxonal spheroids	65743	Autosomal dominant multiple pterygium syndrome
276575	Autosomal dominant hyperinsulinism due to SUR1 deficiency	266	Autosomal dominant limb-girdle muscular dystrophy type 1A	99846	Autosomal dominant myoglobinuria
		264	Autosomal dominant limb-girdle muscular dystrophy type 1B	440354	Autosomal dominant myopia-midfacial retrusion-sensorineural hearing loss-rhizomelic dysplasia syndrome
428	Autosomal dominant hypocalcemia	265	Autosomal dominant limb-girdle muscular dystrophy type 1C	79153	Autosomal dominant nail dysplasia
1810	Autosomal dominant hypohidrotic ectodermal dysplasia	34516	Autosomal dominant limb-girdle muscular dystrophy type 1D	329211	Autosomal dominant neovascular inflammatory vitreoretinopathy
89937	Autosomal dominant hypophosphatemia	34517	Autosomal dominant limb-girdle muscular dystrophy type 1E	98784	Autosomal dominant nocturnal frontal lobe epilepsy
89937	Autosomal dominant hypophosphatemic rickets	55595	Autosomal dominant limb-girdle muscular dystrophy type 1F	178469	Autosomal dominant non-syndromic intellectual disability
457193	Autosomal dominant intellectual disability-craniofacial anomalies-cardiac defects syndrome	55596	Autosomal dominant limb-girdle muscular dystrophy type 1G	90635	Autosomal dominant non-syndromic neurosensory deafness type DFNA
100043	Autosomal dominant intermediate Charcot-Marie-Tooth disease type A	238755	Autosomal dominant limb-girdle muscular dystrophy type 1H	90635	Autosomal dominant non-syndromic neurosensory hearing loss type DFNA
100044	Autosomal dominant intermediate Charcot-Marie-Tooth disease type B	140957	Autosomal dominant macrothrombocytopenia	90635	Autosomal dominant non-syndromic sensorineural deafness type DFNA
100045	Autosomal dominant intermediate Charcot-Marie-Tooth disease type C	88950	Autosomal dominant medullary cystic kidney disease with hyperuricemia	90635	Autosomal dominant non-syndromic sensorineural deafness type DFNA
100046	Autosomal dominant intermediate Charcot-Marie-Tooth disease type D	34149	Autosomal dominant medullary cystic kidney disease with or without hyperuricemia	90635	Autosomal dominant non-syndromic sensorineural hearing loss type DFNA
93114	Autosomal dominant intermediate Charcot-Marie-Tooth disease type E	88949	Autosomal dominant medullary cystic kidney disease without hyperuricemia	93328	Autosomal dominant omodysplasia
352670	Autosomal dominant intermediate Charcot-Marie-Tooth disease type F			306588	Autosomal dominant Opitz BBB/G syndrome
324585	Autosomal dominant intermediate Charcot-Marie-Tooth disease with neuropathic pain	319581	Autosomal dominant mendelian susceptibility to mycobacterial diseases due to partial IFN $\gamma$ R1 deficiency	306588	Autosomal dominant Opitz G/BBB syndrome
90635	Autosomal dominant isolated neurosensory deafness type DFNA	319589	Autosomal dominant mendelian susceptibility to mycobacterial diseases due to partial IFN $\gamma$ R2 deficiency	306588	Autosomal dominant Opitz syndrome
90635	Autosomal dominant isolated neurosensory hearing loss type DFNA			67036	Autosomal dominant optic atrophy and cataract
90635	Autosomal dominant isolated sensorineural deafness type DFNA	319581	Autosomal dominant mendelian susceptibility to mycobacterial diseases due to partial interferon gamma receptor 1 deficiency	→1215	Autosomal dominant optic atrophy and congenital deafness
90635	Autosomal dominant isolated sensorineural hearing loss type DFNA			250932	Autosomal dominant optic atrophy and peripheral neuropathy
93325	Autosomal dominant Kenny-Caffey syndrome	319589	Autosomal dominant mendelian susceptibility to mycobacterial diseases due to partial interferon gamma receptor 2 deficiency	1215	Autosomal dominant optic atrophy plus syndrome
2334	Autosomal dominant keratitis			67036	Autosomal dominant optic atrophy type 3
293936	Autosomal dominant keratoconus with early-onset anterior polar cataracts	2514	Autosomal dominant microcephaly	98673	Autosomal dominant optic atrophy, classic form
503	Autosomal dominant Larsen syndrome	457050	Autosomal dominant mitochondrial myopathy with exercise intolerance	98673	Autosomal dominant optic atrophy, Kjer type
		319581	Autosomal dominant MSMD due to partial IFN $\gamma$ R1 deficiency		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2783	Autosomal dominant osteopetrosis type 1	100988	Autosomal dominant spastic paraplegia type 6	139485	Autosomal recessive ataxia due to coenzyme Q10 deficiency
1798	Autosomal dominant osteosclerosis, Stanescu type	100989	Autosomal dominant spastic paraplegia type 8	247815	Autosomal recessive ataxia due to PEX10 deficiency
2790	Autosomal dominant osteosclerosis, Worth type	100990	Autosomal dominant spastic paraplegia type 9	139485	Autosomal recessive ataxia due to ubiquinone deficiency
1010	Autosomal dominant palmoplantar hyperkeratosis and congenital alopecia	100991	Autosomal dominant spastic paraplegia type 10	88644	Autosomal recessive ataxia, Beauce type
1010	Autosomal dominant palmoplantar keratoderma and congenital alopecia	100993	Autosomal dominant spastic paraplegia type 12	101101	Autosomal recessive axonal Charcot-Marie-Tooth disease type 2B2
		100994	Autosomal dominant spastic paraplegia type 13	101097	Autosomal recessive axonal Charcot-Marie-Tooth disease type 2K
		100998	Autosomal dominant spastic paraplegia type 17	443950	Autosomal recessive axonal Charcot-Marie-Tooth disease type 2T
88924	Autosomal dominant polycystic kidney disease type 1 with tuberous sclerosis	100999	Autosomal dominant spastic paraplegia type 19	90119	Autosomal recessive axonal Charcot-Marie-Tooth disease with acrodystrophy
2924	Autosomal dominant polycystic liver disease	101009	Autosomal dominant spastic paraplegia type 29	98856	Autosomal recessive axonal CMT4C1
1300	Autosomal dominant popliteal pterygium syndrome	101011	Autosomal dominant spastic paraplegia type 31	101102	Autosomal recessive axonal CMT4C2
476119	Autosomal dominant preaxial polydactyly-upperback hypertrichosis syndrome	320365	Autosomal dominant spastic paraplegia type 36	101101	Autosomal recessive axonal CMT4C3
34528	Autosomal dominant primary hypomagnesemia with hypocalciuria	171612	Autosomal dominant spastic paraplegia type 37	101097	Autosomal recessive axonal CMT4C4
2964	Autosomal dominant prognathism	171617	Autosomal dominant spastic paraplegia type 38	324442	Autosomal recessive axonal neuropathy with neuromyotonia
254892	Autosomal dominant progressive external ophthalmoplegia	320355	Autosomal dominant spastic paraplegia type 41	139455	Autosomal recessive bestrophinopathy
88659	Autosomal dominant progressive nephropathy with hypertension	171863	Autosomal dominant spastic paraplegia type 42	448242	Autosomal recessive brachyolmia
314889	Autosomal dominant proximal renal tubular acidosis	444099	Autosomal dominant spastic paraplegia type 73	→3460	Autosomal recessive carpotarsal osteolysis
171871	Autosomal dominant pseudohypoaldosteronism type 1	1797	Autosomal dominant spondylocostal dysostosis	169186	Autosomal recessive centronuclear myopathy
209867	Autosomal dominant rhegmatogenous retinal detachment	1797	Autosomal dominant spondylocostal dysplasia	453521	Autosomal recessive cerebellar ataxia due to CWF19L1 deficiency
3107	Autosomal dominant Robinow syndrome	228169	Autosomal dominant striatal neurodegeneration	352641	Autosomal recessive cerebellar ataxia due to GBA2 deficiency
247511	Autosomal dominant secondary erythrocytosis	98757	Autosomal dominant striatonigral degeneration	412057	Autosomal recessive cerebellar ataxia due to STUB1 deficiency
247511	Autosomal dominant secondary polycythemia	466806	Autosomal dominant thrombocytopenia with platelet secretion defect	88644	Autosomal recessive cerebellar ataxia type 1
98808	Autosomal dominant Segawa syndrome	3357	Autosomal dominant trichoodontoonychodysplasia-syndactyly	139485	Autosomal recessive cerebellar ataxia type 2
486	Autosomal dominant severe congenital neutropenia	3086	Autosomal dominant vitreoretinchoroidopathy	352641	Autosomal recessive cerebellar ataxia with late-onset spasticity
140481	Autosomal dominant slowed nerve conduction velocity	79278	Autosomal erythropoietic protoporphyria	95433	Autosomal recessive cerebellar ataxia-blindness-deafness syndrome
251282	Autosomal dominant spastic ataxia type 1	88919	Autosomal recessive Alport syndrome	352403	Autosomal recessive cerebellar ataxia-cognitive defect syndrome
1182	Autosomal dominant spastic ataxia type 7	1027	Autosomal recessive amelia	404499	Autosomal recessive cerebellar ataxia-epilepsy-intellectual disability syndrome due to KIAA0226 deficiency
100984	Autosomal dominant spastic paraplegia type 3	248	Autosomal recessive anhidrotic ectodermal dysplasia		
100985	Autosomal dominant spastic paraplegia type 4	1116	Autosomal recessive aplasia cutis		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
404493	Autosomal recessive cerebellar ataxia-epilepsy-intellectual disability syndrome due to TUD deficiency	99951	Autosomal recessive congenital hypomyelinating neuropathy	79408	Autosomal recessive dystrophic epidermolysis bullosa, Hallopeau-Siemens type
284282	Autosomal recessive cerebellar ataxia-epilepsy-intellectual disability syndrome due to WWOX deficiency	90349	Autosomal recessive cutis laxa type 1	98855	Autosomal recessive Emery-Dreifuss muscular dystrophy
		357074	Autosomal recessive cutis laxa type 2, classic type	289586	Autosomal recessive exfoliative ichthyosis
284271	Autosomal recessive cerebellar ataxia-psychomotor retardation syndrome	357074	Autosomal recessive cutis laxa type 2, Debré type	1974	Autosomal recessive faciodigitogenital syndrome
		357064	Autosomal recessive cutis laxa type 2, progeroid type	329329	Autosomal recessive frontotemporal pachygyria
363429	Autosomal recessive cerebellar ataxia-pyramidal signs-nystagmus-oculomotor apraxia syndrome	357058	Autosomal recessive cutis laxa type 2A	331226	Autosomal recessive hyper-IgE syndrome due to TYK2 deficiency
95434	Autosomal recessive cerebellar ataxia-saccadic intrusion syndrome	357064	Autosomal recessive cutis laxa type 2B	79644	Autosomal recessive hyperinsulinemic hypoglycemia due to Kir6.2 deficiency
1170	Autosomal recessive cerebelloparenchymal disorder type 3	90349	Autosomal recessive cutis laxa with severe systemic involvement		79643
363969	Autosomal recessive cerebral atrophy	90349	Autosomal recessive cutis laxa, pulmonary emphysema type	79644	Autosomal recessive hyperinsulinism due to Kir6.2 deficiency
466775	Autosomal recessive Charcot Marie Tooth disease type 2X	79500	Autosomal recessive deafness-onychodystrophy syndrome	79643	Autosomal recessive distal osteolysis syndrome
466775	Autosomal recessive Charcot-Marie-Tooth disease type 2 due to SPG11 mutation	2776	Autosomal recessive distal renal tubular acidosis		79643
324442	Autosomal recessive Charcot-Marie-Tooth disease type 2 with neuromyotonia	→402041	Autosomal recessive distal renal tubular acidosis with deafness	248	Autosomal recessive hypohidrotic ectodermal dysplasia
		→402041	Autosomal recessive distal renal tubular acidosis with hearing loss	289176	Autosomal recessive hypophosphatemic rickets
98856	Autosomal recessive Charcot-Marie-Tooth disease type 2B1	→402041	Autosomal recessive distal renal tubular acidosis without deafness	300547	Autosomal recessive infantile hypercalcemia
101097	Autosomal recessive Charcot-Marie-Tooth disease with hoarseness	→402041	Autosomal recessive distal renal tubular acidosis without hearing loss	93591	Autosomal recessive infantile nephronophthisis
90118	Autosomal recessive Charcot-Marie-Tooth disease, Ouvrier type	402041	Autosomal recessive distal RTA	93591	Autosomal recessive infantile NPHP
90119	Autosomal recessive Charcot-Marie-Tooth type 2 with acrodystrophy	→402041	Autosomal recessive distal RTA with deafness	352530	Autosomal recessive intellectual disability due to TRAPPC9 deficiency
293603	Autosomal recessive CHED	98920	Autosomal recessive distal spinal muscular atrophy type 1	217055	Autosomal recessive intermediate Charcot-Marie-Tooth disease type A
2518	Autosomal recessive chorioretinopathy-microcephaly syndrome		139552	Autosomal recessive distal spinal muscular atrophy type 2	254334
447760	Autosomal recessive complex spastic paraplegia type 9B	139547	Autosomal recessive distal spinal muscular atrophy type 3	369867	Autosomal recessive intermediate Charcot-Marie-Tooth disease type C
363432	Autosomal recessive congenital cerebellar ataxia due to GRID2 deficiency	206580	Autosomal recessive distal spinal muscular atrophy type 4	435998	Autosomal recessive intermediate Charcot-Marie-Tooth disease type D
363432	Autosomal recessive congenital cerebellar ataxia due to ionotropic glutamate receptor delta-2 subunit deficiency	314485	Autosomal recessive distal spinal muscular atrophy type 5	210110	Autosomal recessive intermediate osteopetrosis
		101150	Autosomal recessive dopa-responsive dystonia	90636	Autosomal recessive isolated neurosensory deafness type DFNB
324262	Autosomal recessive congenital cerebellar ataxia due to metabotropic glutamate receptor 1 deficiency	79408	Autosomal recessive dystrophic epidermolysis bullosa generalisata gravis	98676	Autosomal recessive isolated optic atrophy
		89842	Autosomal recessive dystrophic epidermolysis bullosa generalisata mitis	90636	Autosomal recessive isolated sensorineural deafness type DFNB
324262	Autosomal recessive congenital cerebellar ataxia due to MGLUR1 deficiency	89842	Autosomal recessive dystrophic epidermolysis bullosa, generalized other	93324	Autosomal recessive Kenny-Caffey syndrome
293603	Autosomal recessive congenital hereditary endothelial dystrophy				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
1842	Autosomal recessive lethal chondrodysplasia, round femoral inferior epiphysis type	363623	Autosomal recessive limb-girdle muscular dystrophy type 2T	319574	Autosomal recessive MSMD due to partial interferon gamma receptor 2 deficiency
33108	Autosomal recessive lethal multiple pterygium syndrome	352479	Autosomal recessive limb-girdle muscular dystrophy type 2U	93307	Autosomal recessive multiple epiphyseal dysplasia
314572	Autosomal recessive leukoencephalopathy-ischemic stroke-retinitis pigmentosa syndrome	466801	Autosomal recessive limb-girdle muscular dystrophy type 2W	2990	Autosomal recessive multiple pterygium syndrome
		476084	Autosomal recessive limb-girdle muscular dystrophy type 2X	424261	Autosomal recessive muscular dystrophy due to LAP1B deficiency
		424261	Autosomal recessive limb-girdle muscular dystrophy type 2Y	424261	Autosomal recessive muscular dystrophy due to Torsin-1A-interacting protein 1 deficiency
363543	Autosomal recessive limb-girdle muscular dystrophy due to desmin deficiency	480682	Autosomal recessive limb-girdle muscular dystrophy type 2Z	319332	Autosomal recessive myogenic AMC
352479	Autosomal recessive limb-girdle muscular dystrophy due to ISPD deficiency	476084	Autosomal recessive limb-girdle muscular dystrophy-cardiac arrhythmia syndrome	319332	Autosomal recessive myogenic arthrogryposis multiplex congenita
254361	Autosomal recessive limb-girdle muscular dystrophy due to plectin deficiency	206580	Autosomal recessive lower motor neuron disease with childhood onset	280654	Autosomal recessive nail dysplasia
267	Autosomal recessive limb-girdle muscular dystrophy type 2A	238505	Autosomal recessive lymphoproliferative disease	2990	Autosomal recessive non-lethal multiple pterygium syndrome
268	Autosomal recessive limb-girdle muscular dystrophy type 2B	667	Autosomal recessive malignant osteopetrosis	88616	Autosomal recessive non-syndromic intellectual disability
353	Autosomal recessive limb-girdle muscular dystrophy type 2C	477857	Autosomal recessive mendelian susceptibility to mycobacterial diseases due to complete RORgamma receptor deficiency	90636	Autosomal recessive non-syndromic neurosensory deafness type DFNB
62	Autosomal recessive limb-girdle muscular dystrophy type 2D			90636	Autosomal recessive non-syndromic sensorineural deafness type DFNB
119	Autosomal recessive limb-girdle muscular dystrophy type 2E			93329	Autosomal recessive omodysplasia
219	Autosomal recessive limb-girdle muscular dystrophy type 2F	319569	Autosomal recessive mendelian susceptibility to mycobacterial diseases due to partial IFNgammaR1 deficiency	67047	Autosomal recessive optic atrophy plus syndrome
34514	Autosomal recessive limb-girdle muscular dystrophy type 2G	319574	Autosomal recessive mendelian susceptibility to mycobacterial diseases due to partial IFNgammaR2 deficiency	67047	Autosomal recessive optic atrophy type 3
1878	Autosomal recessive limb-girdle muscular dystrophy type 2H	319569	Autosomal recessive mendelian susceptibility to mycobacterial diseases due to partial interferon gamma receptor 1 deficiency	227976	Autosomal recessive optic atrophy, OPA7 type
34515	Autosomal recessive limb-girdle muscular dystrophy type 2I			178389	Autosomal recessive osteoclast-poor osteopetrosis with hypogammaglobulinemia
140922	Autosomal recessive limb-girdle muscular dystrophy type 2J			178389	Autosomal recessive osteopetrosis type 7
86812	Autosomal recessive limb-girdle muscular dystrophy type 2K	319574	Autosomal recessive mendelian susceptibility to mycobacterial diseases due to partial interferon gamma receptor 2 deficiency	1366	Autosomal recessive palmoplantar hyperkeratosis and congenital alopecia
206549	Autosomal recessive limb-girdle muscular dystrophy type 2L	175	Autosomal recessive metaphyseal chondrodysplasia	1366	Autosomal recessive palmoplantar keratoderma and congenital alopecia
206554	Autosomal recessive limb-girdle muscular dystrophy type 2M	621	Autosomal recessive methemoglobinemia	731	Autosomal recessive polycystic kidney disease
206559	Autosomal recessive limb-girdle muscular dystrophy type 2N	477857	Autosomal recessive MSMD due to complete RORgamma receptor deficiency	1234	Autosomal recessive popliteal pterygium syndrome
206564	Autosomal recessive limb-girdle muscular dystrophy type 2O	319569	Autosomal recessive MSMD due to partial IFNgammaR1 deficiency	88628	Autosomal recessive posterior column ataxia and retinitis pigmentosa
280333	Autosomal recessive limb-girdle muscular dystrophy type 2P	319574	Autosomal recessive MSMD due to partial IFNgammaR2 deficiency	477857	Autosomal recessive primary immunodeficiency due to RORC mutation
363543	Autosomal recessive limb-girdle muscular dystrophy type 2R	319569	Autosomal recessive MSMD due to partial interferon gamma receptor 1 deficiency	437552	Autosomal recessive primary immunodeficiency with defective spontaneous natural killer cell cytotoxicity
369840	Autosomal recessive limb-girdle muscular dystrophy type 2S				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
437552	Autosomal recessive primary immunodeficiency with defective spontaneous NK cell cytotoxicity	314603	Autosomal recessive spastic ataxia with leukoencephalopathy	431329	Autosomal recessive spastic paraplegia type 57
2512	Autosomal recessive primary microcephaly	254343	Autosomal recessive spastic ataxia-optic atrophy-dysarthria syndrome	397946	Autosomal recessive spastic paraplegia type 58
254886	Autosomal recessive progressive external ophthalmoplegia	100986	Autosomal recessive spastic paraplegia type 5A	401795	Autosomal recessive spastic paraplegia type 59
93607	Autosomal recessive proximal renal tubular acidosis	2822	Autosomal recessive spastic paraplegia type 11	401800	Autosomal recessive spastic paraplegia type 60
171876	Autosomal recessive pseudohypoaldosteronism type 1	100995	Autosomal recessive spastic paraplegia type 14	401780	Autosomal recessive spastic paraplegia type 61
1507	Autosomal recessive Robinow syndrome	100996	Autosomal recessive spastic paraplegia type 15	401785	Autosomal recessive spastic paraplegia type 62
247378	Autosomal recessive secondary erythrocytosis not associated with VHL gene	209951	Autosomal recessive spastic paraplegia type 18	401805	Autosomal recessive spastic paraplegia type 63
247378	Autosomal recessive secondary erythrocytosis, non-Chuvash type	101000	Autosomal recessive spastic paraplegia type 20	401810	Autosomal recessive spastic paraplegia type 64
247378	Autosomal recessive secondary polycythemia not associated with VHL gene	101001	Autosomal recessive spastic paraplegia type 21	320396	Autosomal recessive spastic paraplegia type 65
247378	Autosomal recessive secondary polycythemia, non-Chuvash type	101003	Autosomal recessive spastic paraplegia type 23	401815	Autosomal recessive spastic paraplegia type 66
247378	Autosomal recessive secondary polycythemia, non-Chuvash type	101004	Autosomal recessive spastic paraplegia type 24	401820	Autosomal recessive spastic paraplegia type 67
101150	Autosomal recessive Segawa syndrome	101005	Autosomal recessive spastic paraplegia type 25	401825	Autosomal recessive spastic paraplegia type 68
970	Autosomal recessive sensory radicular neuropathy	101006	Autosomal recessive spastic paraplegia type 26	401830	Autosomal recessive spastic paraplegia type 69
70594	Autosomal recessive sepiapterin reductase-deficient DRD	101007	Autosomal recessive spastic paraplegia type 27	401835	Autosomal recessive spastic paraplegia type 70
420702	Autosomal recessive severe congenital neutropenia due to CSF3R deficiency	101008	Autosomal recessive spastic paraplegia type 28	401840	Autosomal recessive spastic paraplegia type 71
420699	Autosomal recessive severe congenital neutropenia due to CXCR2 deficiency	171622	Autosomal recessive spastic paraplegia type 32	468661	Autosomal recessive spastic paraplegia type 74
331176	Autosomal recessive severe congenital neutropenia due to G6PC3 deficiency	171629	Autosomal recessive spastic paraplegia type 35	459056	Autosomal recessive spastic paraplegia type 75
423384	Autosomal recessive severe congenital neutropenia due to JAGN1 deficiency	139480	Autosomal recessive spastic paraplegia type 39	466722	Autosomal recessive spastic paraplegia type 77
260305	Autosomal recessive sideroblastic anemia	320370	Autosomal recessive spastic paraplegia type 43	101005	Autosomal recessive spastic paraplegia-disc herniation syndrome
98	Autosomal recessive spastic ataxia of Charlevoix-Saguenay	320401	Autosomal recessive spastic paraplegia type 44	98920	Autosomal recessive spinal muscular atrophy with respiratory distress
397946	Autosomal recessive spastic ataxia type 2	320396	Autosomal recessive spastic paraplegia type 45	1170	Autosomal recessive spinocerebellar ataxia type 2
314603	Autosomal recessive spastic ataxia type 3	320391	Autosomal recessive spastic paraplegia type 46	284332	Autosomal recessive spinocerebellar ataxia type 6
254343	Autosomal recessive spastic ataxia type 4	306511	Autosomal recessive spastic paraplegia type 48	284324	Autosomal recessive spinocerebellar ataxia type 7
313772	Autosomal recessive spastic ataxia type 5	320385	Autosomal recessive spastic paraplegia type 49	139485	Autosomal recessive spinocerebellar ataxia type 9
98	Autosomal recessive spastic ataxia type 6	319199	Autosomal recessive spastic paraplegia type 53	284289	Autosomal recessive spinocerebellar ataxia type 10
		320380	Autosomal recessive spastic paraplegia type 54	284271	Autosomal recessive spinocerebellar ataxia type 11
		320375	Autosomal recessive spastic paraplegia type 55	284282	Autosomal recessive spinocerebellar ataxia type 12
		320411	Autosomal recessive spastic paraplegia type 56		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
324262	Autosomal recessive spinocerebellar ataxia type 13	209004	Axonal polyneuropathy associated with IgG/IgM/IgA monoclonal gammopathy	1227	Bangstad syndrome
352403	Autosomal recessive spinocerebellar ataxia type 14		1435	Ayazi syndrome	130
404499	Autosomal recessive spinocerebellar ataxia type 15	477668	Aymé-Gripp syndrome	1228	Banki syndrome
397709	Autosomal recessive spinocerebellar ataxia type 20	284454	AZOOR	109	Bannayan-Riley-Ruvalcaba syndrome
		→399805	Azoospermia due to maturation arrest	139507	Bantu siderosis
466794	Autosomal recessive spinocerebellar ataxia type 21	→399805	Azoospermia due to meiosis defect	289539	BAP1-related tumor predisposition syndrome
2311	Autosomal recessive spondylocostal dysostosis	3471	Azoospermia-sinopulmonary infections syndrome	1229	Baraitser-Brett-Piesowicz syndrome
		98757	Azorean disease of the nervous system	2753	Baraitser-Burn syndrome
401979	Autosomal recessive spondylometaphyseal dysplasia, Mégarbané type	99121	Azygos continuation of the inferior caval vein	1229	Baraitser-Reardon syndrome
250984	Autosomal recessive Stickler syndrome	99121	Azygos continuation of the inferior vena cava	2995	Baraitser-Winter cerebrofrontofacial syndrome
		99121	Azygos continuation of the IVC	2237	Barakat syndrome
280365	Autosomal semi-dominant severe lipodystrophic laminopathy	79332	B4GALT1-CDG	1231	Barber-Say syndrome
300345	Autosomal SLE	75496	B4GALT7-CDG	110	Bardet-Biedl syndrome
101010	Autosomal spastic paraplegia type 30	99860	B-ALL	34592	Bare lymphocyte syndrome type 1
401849	Autosomal spastic paraplegia type 72	67038	B-cell chronic lymphocytic leukemia	572	Bare lymphocyte syndrome type 2
		67038	B-cell chronic lymphoid leukemia	3317	Barnes syndrome
300345	Autosomal systemic lupus erythematosus	464336	B-cell expansion with NF-κB and T-cell anergy disease	443084	Baroreflex failure
168629	Autosomal thrombocytopenia with normal platelets	86852	B-cell prolymphocytic leukemia	79087	Barraquer-Simons syndrome
		67038	B-CLL	2698	Bart-Pumphrey syndrome
352490	AUTS2 syndrome	404560	B-K mole syndrome	111	Barth syndrome
96	AVED	86852	B-PLL	64692	Bartonellosis due to Bartonella bacilliformis infection
98963	Avellino corneal dystrophy	108	Babesiosis	50839	Bartonellosis due to Bartonella henselae infection
454836	Avian influenza	206994	Bacterial myositis	64694	Bartonellosis due to Bartonella quintana infection
99000	AVMD	36234	Bacterial toxic-shock syndrome	1234	Bartsocas-Papas syndrome
58	AxD	36234	Bacterial TSS	112	Bartter syndrome
363717	AxD type I	69736	BADI	93605	Bartter syndrome type 3
363722	AxD type II	86814	BAFME	89938	Bartter syndrome type 4
98978	Axenfeld anomaly	2819	Bahemuka-Brown syndrome	263417	Bartter syndrome type 5
782	Axenfeld syndrome	352577	Bainbridge-Roppers syndrome	93605	Bartter syndrome type III
782	Axenfeld-Rieger syndrome	1658	Baird syndrome	89938	Bartter syndrome type IV
1834	Axial mesodermal dysplasia spectrum	139471	Bakrania-Ragge syndrome	263417	Bartter syndrome with hypocalcemia
		1223	Balantidiasis	93604	Bartter syndrome, furosemide type
2777	Axial osteosclerosis	1223	Balantidiasis	93604	Bartter syndrome, furosemide-amiloride type
168549	Axial spondylometaphyseal dysplasia	139450	Balikova-Vermeesch syndrome	377	Basal cell nevus syndrome
		363746	Balint syndrome	268829	Basal encephalocele
401911	AXIN2-related AFAP	363746	Balint-Holmes syndrome	→1658	Basan syndrome
401911	AXIN2-related attenuated familial adenomatous polyposis	93395	Ballard syndrome	79301	BASD1
		1225	Baller-Gerold syndrome	79303	BASD2
401911	AXIN2-related attenuated familial polyposis coli	66529	Ballooning cardiomyopathy	79302	BASD3
		228165	Baló concentric sclerosis	79095	BASD4
401911	AXIN2-related attenuated FAP	634	Bamboo hair syndrome	50810	Basel-Vanagaite-Sirota syndrome
101102	Axonal Charcot-Marie-Tooth disease with pyramidal involvement	1226	Bamforth syndrome	244283	BASM syndrome
		1226	Bamforth-Lazarus syndrome	14	Bassen-Kornzweig disease
457205	Axonal neuropathy-optic atrophy-cognitive deficit syndrome	98955	Band-shaped and whorled microcystic dystrophy of the corneal epithelium	1875	Bassoe syndrome
				100976	Bathing suit ichthyosis

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
1948	Battaglia-Neri syndrome	247203	Bellini duct carcinoma	238624	Benign intracranial hypertension
79264	Batten disease	1240	Bellini syndrome	285	Benign joint hypermobility syndrome
→1071	Baughman syndrome	1492	Ben Ari-Shuper-Mimouni syndrome	464359	Benign metanephric tumour
166113	Bazex syndrome	100978	Benallegue-Lacete syndrome	168816	Benign multicystic peritoneal mesothelioma
113	Bazex-Dupré-Christol syndrome	1241	Bencze syndrome	86909	Benign myoclonic epilepsy of infancy
65284	BBGD	86814	Benign adult familial myoclonic epilepsy	86909	Benign myoclonus epilepsy of infancy
363444	BBIS	86814	Benign adult familial myoclonus epilepsy	140927	Benign neonatal-infantile epilepsy
110	BBS	610	Benign autosomal dominant myopathy	25968	Benign occipital epilepsy
41751	BCD	157997	Benign cephalic histiocytosis	342	Benign paroxysmal peritonitis
1997	BCD syndrome	98816	Benign childhood occipital epilepsy, Gastaut type	1179	Benign paroxysmal tonic upgaze of childhood with ataxia
312	BCIE	98815	Benign childhood occipital epilepsy, Panayiotopoulos type	71518	Benign paroxysmal torticollis of infancy
511	BCKD deficiency	2841	Benign chronic familial pemphigus of Hailey-Hailey	166299	Benign partial epilepsy of infancy with complex partial seizures
511	BCKDH deficiency	251287	Benign concentric annular macular dystrophy	166302	Benign partial epilepsy with secondarily generalized seizures in infancy
247203	BDC	440233	Benign congenital sixth cranial nerve palsy	65682	Benign recurrent intrahepatic cholestasis
113	BDCS	254864	Benign COX deficiency	99960	Benign recurrent intrahepatic cholestasis type 1
115	Beals syndrome	180284	Benign ductal tumor of breast	99961	Benign recurrent intrahepatic cholestasis type 2
115	Beals-Hecht syndrome	1945	Benign epilepsy of childhood with centrotemporal spikes	342	Benign recurrent polyserositis
1059	Bean syndrome	276148	Benign epithelial tumor of salivary glands	1945	Benign rolandic epilepsy
1555	Beare-Stevenson cutis gyrata syndrome	71269	Benign exophthalmos syndrome	324581	Benign Samaritan congenital myopathy
363444	Beaulieu-Boycott-Innes syndrome	1429	Benign familial chorea	252164	Benign schwannoma
98895	Becker dystrophinopathy	1945	Benign familial epilepsy of childhood with rolandic spikes	180237	Benign tumor of fallopian tubes
98895	Becker muscular dystrophy	306	Benign familial infantile convulsions	2198	Bennion-Patterson syndrome
64755	Becker nevus syndrome	306	Benign familial infantile epilepsy	54247	Benson syndrome
116	Beckwith-Wiedemann syndrome	306	Benign familial infantile seizures	464336	BENTA disease
231127	Beckwith-Wiedemann syndrome due to 11p15 microdeletion	163717	Benign familial mesial temporal lobe epilepsy	528	Berardinelli-Seip syndrome
96076	Beckwith-Wiedemann syndrome due to 11p15 microduplication	1949	Benign familial neonatal convulsions	171839	Berant syndrome
231130	Beckwith-Wiedemann syndrome due to 11p15 translocation/inversion	1949	Benign familial neonatal epilepsy	528	Berardinelli-Seip congenital lipodystrophy
231120	Beckwith-Wiedemann syndrome due to CDKN1C mutation	1949	Benign familial neonatal seizures	2241	Berdon syndrome
231117	Beckwith-Wiedemann syndrome due to imprinting defect of 11p15	140927	Benign familial neonatal-infantile seizures	647	Berlin breakage syndrome
238613	Beckwith-Wiedemann syndrome due to NSD1 mutation	209973	Benign familial nocturnal alternating hemiplegia in childhood	274	Bernard-Soulier syndrome
96193	Beckwith-Wiedemann syndrome due to paternal uniparental disomy of chromosome 11	209973	Benign familial nocturnal alternating hemiplegia of childhood	178528	Berti lymphoma
1945	BE CRS	65684	Benign focal amyotrophy	133	Berylliosis
1945	BE CTS	1544	Benign focal seizures of adolescence	71269	BES
2572	Bedouin spastic ataxia syndrome	64545	Benign idiopathic neonatal seizures	797	Besnier-Boeck-Schaumann disease
322	BEEC	166308	Benign infantile focal epilepsy with midline spikes and wave during sleep	321	Bessel-Hagen disease
1237	Beemer-Ertbruggen syndrome	166305	Benign infantile seizures associated with mild gastroenteritis	1243	Best disease
275864	Behavioral variant of frontotemporal dementia			1243	Best macular dystrophy
1239	Behr syndrome			1243	Best vitelliform macular dystrophy
2705	Behrens-Baumann-Vogel syndrome			79332	Beta-1,4-galactosyltransferase deficiency
117	Behçet disease			65287	Beta-alanine synthase deficiency
476102	Behçet-like disease due to HA20				
476102	Behçet-like disease due to haploinsufficiency of A20				
247203	Bellini carcinoma				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
309310	Beta-D-galactosidase deficiency	295177	Bifid halluces, bilateral	386	Biliary hamartoma
354	Beta-galactosidase-1 deficiency	295175	Bifid halluces, unilateral	→2697	Biliary tract malformation-renal failure syndrome
584	Beta-glucuronidase deficiency	295006	Bifid hallux	98836	Bilineal acute leukemia
134	Beta-ketothiolase deficiency	295177	Bifid hallux, bilateral	415286	Bilirubin encephalopathy
118	Beta-mannosidase deficiency	295175	Bifid hallux, unilateral	205	Bilirubin uridinediphosphate glucuronosyltransferase deficiency
118	Beta-mannosidosis	2695	Bifid nose	79234	Bilirubin uridinediphosphate glucuronosyltransferase deficiency type 1
329284	Beta-propeller protein-associated neurodegeneration	217266	Bifid nose with or without anorectal and renal anomalies	79235	Bilirubin uridinediphosphate glucuronosyltransferase deficiency type 2
119	Beta-sarcoglycanopathy	99771	Bifid uvula	205	Bilirubin-UGT deficiency
848	Beta-thalassemia	99771	Bifidity of the uvula	79234	Bilirubin-UGT deficiency type 1
231222	Beta-thalassemia intermedia	300	Bifunctional enzyme deficiency	79235	Bilirubin-UGT deficiency type 2
231214	Beta-thalassemia major	637	Bilateral acoustic neurofibromatosis	205	Bilirubin-UGT deficiency
→33364	Beta-thalassemia-trichothiodystrophy syndrome	69736	Bilateral acute depigmentation of the iris	79234	Bilirubin-UGT deficiency type 1
231393	Beta-thalassemia-X-linked thrombocytopenia syndrome	319205	Bilateral adrenal hemorrhage	79235	Bilirubin-UGT deficiency type 2
65287	Beta-ureidopropionase deficiency	325124	Bilateral anorchia	1799	Billard-Toutain-Maheut syndrome
610	Bethlem myopathy	2048	Bilateral anterior opercular syndrome	1248	Binder syndrome
2114	Beukes familial hip dysplasia	1229	Bilateral band-like calcification with polymicrogyria	3304	Bindewald-Ulmer-Müller syndrome
2114	BFHD	208444	Bilateral frontal polymicrogyria	65284	Biotin-responsive basal ganglia disease
306	BFIE	101070	Bilateral frontoparietal polymicrogyria	65284	Biotin-thiamine-responsive basal ganglia disease
306	BFIS	208447	Bilateral generalized polymicrogyria	79241	Biotinidase deficiency
127	BFLS	93311	Bilateral hereditary micro-epiphyseal dysplasia	54247	Biparietal Alzheimer disease
140927	BFNIS	438117	Bilateral hip and radial head dislocations-short stature-scoliosis-carpal coalitions-pes cavus-facial dysmorphism syndrome	364198	Bipartite talus
1949	BFNS	319205	Bilateral massive adrenal hemorrhage	99908	Bird fancier lung
293284	BH4-responsive HPA/PKU	97364	Bilateral MC DK	2617	Bird headed-dwarfism, Montreal type
293284	BH4-responsive hyperphenylalaninemia/phenylketonuria	140963	Bilateral microtia-deafness-cleft palate syndrome	179	Birdshot chorioretinitis
93311	BHMED	97364	Bilateral multicystic dysplastic kidney	179	Birdshot chorioretinopathy
98964	Biber-Haab-Dimmer dystrophy	97364	Bilateral multicystic renal dysplasia	179	Birdshot retinochoroiditis
180086	Bicervical bicornuate uterus	208441	Bilateral parasagittal parieto-occipital polymicrogyria	179	Birdshot retinochoroidopathy
180106	Bicervical bicornuate uterus and blind hemivagina	98889	Bilateral perisylvian polymicrogyria	122	Birt-Hogg-Dubé syndrome
180111	Bicervical bicornuate uterus with patent cervix and vagina	268940	Bilateral polymicrogyria	79133	Bitemporal aplasia cutis congenita
2088	Bickel-Fanconi glycogenosis	295150	Bilateral PPD2	2213	Bixler-Christian-Gorlin syndrome
2182	Bickers-Adams syndrome	1980	Bilateral striopallidodentate calcinosis	285	BJHS
79138	Bickerstaff brainstem encephalitis	276066	Bile acid CoA ligase deficiency and defective amidation	123	Björnstad syndrome
3286	Bidirectional tachycardia induced by catecholamine	70567	Bile duct cancer	124	Blackfan-Diamond anemia
→33364	BIDS syndrome	1276	Bilginturan brachydactyly	93930	Bladder exstrophy
1246	Biemond syndrome	1276	Bilginturan syndrome	322	Bladder exstrophy-epispadias-cloacal extrophy complex
141333	Biemond syndrome type 2	1247	Bilharziasis	37202	Bladder pain syndrome
41751	Bietti crystalline corneoretinal dystrophy	30391	Biliary atresia	98922	Blake pouch cyst
41751	Bietti crystalline dystrophy	244283	Biliary atresia with splenic malformation syndrome	254379	Blaschkoid lichen planus
41751	Bietti crystalline retinopathy	424982	Biliary cystadenocarcinoma	254379	Blaschkoid LP
1986	Bifid femur-monodactylous ectrodactyly syndrome			86870	Blastic NK-cell lymphoma
295006	Bifid great toes			86870	Blastic plasmacytoid dendritic cell neoplasm
295177	Bifid great toes, bilateral			1834	Blastogenesis defect
295175	Bifid great toes, unilateral			90340	Blau syndrome
295006	Bifid halluces			50945	BLC
				1229	BLC-PMG
				73271	Bleeding diathesis due to a collagen receptor defect

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
98885	Bleeding diathesis due to glycoprotein VI deficiency	1256	Blepharophimosis-radioulnar synostosis syndrome	1842	Bone dysplasia, lethal Holmgren type
98886	Bleeding diathesis due to integrin alpha2-beta1 deficiency	1968	Blepharophimosis-telecanthus-microstomia syndrome	85182	Bone dysplasia-medullary fibrosarcoma syndrome
220443	Bleeding diathesis due to thromboxane synthesis deficiency	1258	Blepharoptosis-cleft palate-ectrodactyly-dental anomalies syndrome	300284	Bone fragility-contractures-arterial rupture-deafness syndrome
420566	Bleeding disorder due to calcium- and DAG-regulated guanine exchange factor-1 deficiency	1259	Blepharoptosis-myopia-ectopia lentis syndrome	2050	Bone fragility-craniosynostosis-proptosis-hydrocephalus syndrome
420566	Bleeding disorder due to Ca/DAG-GEFI deficiency	93964	Blepharospasm-romandibular dystonia syndrome	2934	Bonneau syndrome
1997	Blepharo-cheilo-odontic syndrome	171844	Blepharospasm-romandibular dystonia syndrome	163	Bonneau-Beaumont syndrome
1253	Blepharochalasis-double lip syndrome		464	Blindness-scoliosis-arachnodactyly syndrome	2941
1997	Blepharocheilodontic syndrome	464	Bloch-Siemens syndrome	1261	Bonnemann-Meinecke-Reich syndrome
→2353	Blepharofacioskeletal syndrome	50945	Bloch-Sulzberger syndrome	53719	Bonnet-Dechaume-Blanc syndrome
1252	Blepharonasofacial malformation syndrome	50945	Bloomstrand chondrodysplasia	1262	Böök syndrome
2728	Blepharophimosis syndrome, Ohdo type	50945	Bloomstrand lethal chondrodysplasia	1263	Boomerang dysplasia
126	Blepharophimosis types 1 and 2	125	Bloom syndrome	1303	BOOP
261572	Blepharophimosis types 1 and 2 due to a point mutation	2768	Blount disease	1933	Booth-Haworth-Dilling syndrome
261579	Blepharophimosis types 1 and 2 due to copy number variations	88629	Blue colour blindness	107	BOR syndrome
261559	Blepharophimosis-epicanthus inversus-ptosis due to 3q23 rearrangement syndrome	16	Blue cone monochromacy	206473	Borderline epithelial tumor of ovary
261579	Blepharophimosis-epicanthus inversus-ptosis due to a CNV	16	Blue cone monochromatism	206473	Borderline ovarian epithelial tumor
261572	Blepharophimosis-epicanthus inversus-ptosis due to a point mutation syndrome	94086	Blue diaper syndrome	127	Borjeson-Forssman-Lehmann syndrome
261579	Blepharophimosis-epicanthus inversus-ptosis due to copy number variations	1059	Blue rubber bleb nevus	1264	Bork syndrome
126	Blepharophimosis-epicanthus inversus-ptosis syndrome	98989	Blue-dot cataract	90001	Bornholm eye disease
329255	Blepharophimosis-intellectual disability syndrome due to UBE3B deficiency	319205	BMAH	36273	Borrmann gastric cancer type 4
293725	Blepharophimosis-intellectual disability syndrome type V	1243	BMD	97297	BOS syndrome
293707	Blepharophimosis-intellectual disability syndrome, Maat-Kievit-Brunner type	98895	BMD	69737	Bosley-Salih-Alorainy syndrome
293707	Blepharophimosis-intellectual disability syndrome, MKB type	293725	BMRS type V	2250	Bosma-Henkin-Christiansen syndrome
2728	Blepharophimosis-intellectual disability syndrome, Ohdo type	293707	BMRS, Maat-Kievit-Brunner type	85128	Bothnia retinal dystrophy
3047	Blepharophimosis-intellectual disability syndrome, SBBYS type	293707	BMRS, MKB type	128	Bothriocephalosis
293725	Blepharophimosis-intellectual disability syndrome, Verloes type	2728	BMRS, Ohdo type	1267	Botulism
2057	Blepharophimosis-ptosis-esotropia-syndactyly-short stature syndrome	293725	BMRS, Verloes type	1180	Boucher-Neuhäuser syndrome
		353253	BMS	805	Bourneville syndrome
		217266	BNAR syndrome	83313	Boutonneuse fever
		50945	BOCD	→912	Bowen syndrome
		217008	Bockenheimer syndrome	1270	Bowen syndrome, Hutterite type
		1292	BOD syndrome	1270	Bowen-Conradi syndrome
		2724	Boder syndrome	97353	Boxer's dementia
		48686	Body cavity-based lymphoma	50814	Boydjiev-Jabs syndrome
		91135	Body skin hyperlaxity due to vitamin K-dependent coagulation factor deficiency	2680	Boylan-Dew syndrome
		797	Boeck sarcoid	329284	BPAN
		797	Boeck's sarcoid	70589	BPD
		1297	BOFS	86870	BPDCN
		97297	Bohring syndrome	2901	Brachial plexus neuritis
		97297	Bohring-Opitz syndrome	199	Brachmann-de Lange syndrome
		84081	Boichis disease	1519	Brachycephalofrontonasal dysplasia
		401874	BOLA3 deficiency	1272	Brachycephaly-deafness-cataract-intellectual disability syndrome
		319229	Bolivian hemorrhagic fever	2619	Brachydactylous dwarfism, Mseleni type
		1844	Bone dysplasia, Azouz type	294996	Brachydactyly of fingers
				295130	Brachydactyly of fingers, bilateral

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
295128	Brachydactyly of fingers, unilateral	93302	Brachyolmia, Maroteaux type	→33364	Brittle hair-mental deficiency syndrome
294998	Brachydactyly of toes	2899	Brachyolmia-amelogenesis imperfecta syndrome	783	Broad thumb-hallux syndrome
295134	Brachydactyly of toes, bilateral	79345	Brachytelephalangic chondrodysplasia punctata	783	Broad thumbs-halluces syndrome
295132	Brachydactyly of toes, unilateral	1295	Brachytelephalangy-dysmorphism-Kallmann syndrome	412	Broad-betalipoproteinemia
93388	Brachydactyly type A1	441	Bradbury-Eggleston syndrome	53347	Brody myopathy
93396	Brachydactyly type A2	52047	Braddock syndrome	66529	Broken heart syndrome
93394	Brachydactyly type A4	3323	Braddock-Carey syndrome	97287	Bronchial endocrine tumor
93389	Brachydactyly type A5	1538	Braddock-Jones-Superneau syndrome	→3471	Bronchiectasis-oligospermia syndrome
93382	Brachydactyly type A6	75374	Bradyopsia	1302	Bronchiolitis obliterans organizing pneumonia
93397	Brachydactyly type A7	178506	Brain calcification, Rajab type	1303	Bronchiolitis obliterans with obstructive pulmonary disease
93383	Brachydactyly type B	168598	Brain demyelination due to methionine adenosyltransferase deficiency	2357	Bronchogenic cyst
140908	Brachydactyly type B2	352649	Brain dopamine-serotonin vesicular transport disease	70589	Bronchopulmonary dysplasia
93384	Brachydactyly type C	75389	Brain malformation-congenital heart disease-postaxial polydactyly syndrome	1116	Bronspiegel-Zelnick syndrome
93387	Brachydactyly type E	467166	Brain stem asymmetry-superior cerebellar and basal ganglia dysplasia syndrome	99829	Bronze John
1276	Brachydactyly type E, with short stature and hypertension	209905	Brain-lung-thyroid syndrome	79493	Brooke-Spiegler syndrome
93395	Brachydactyly types B and E combined	255182	Branched chain alpha-ketoacid dehydrogenase complex deficiency	97229	Brown-Vialetto-van Laere syndrome
93388	Brachydactyly, Farabee type	511	Branched-chain 2-ketoacid dehydrogenase deficiency	109	BRRS
2946	Brachydactyly, long thumb type	1296	Branchial dysplasia-intellectual disability-inguinal hernia syndrome	2353	BRSS
93396	Brachydactyly, Mohr-Wriedt type	1297	Branchio-oculo-facial syndrome	1304	Brucellosis
93397	Brachydactyly, Smorgasbord type	50815	Branchiogenic deafness syndrome	2771	Bruck syndrome
93394	Brachydactyly, Temtamy type	52429	Branchiootic syndrome	130	Brugada syndrome
1276	Brachydactyly-arterial hypertension syndrome	107	Branchiootorenal syndrome	3057	Brunner syndrome
1275	Brachydactyly-elbow wrist dysplasia syndrome	1299	Branchioskeletogenital syndrome	1305	Brunner-Winter syndrome
1001	Brachydactyly-intellectual disability syndrome	79133	Brauer syndrome	391641	Brunner-Winter syndrome type 1
1275	Brachydactyly-joint dysplasia syndrome	2669	Braun-Bayer syndrome	391646	Brunner-Winter syndrome type 2
2946	Brachydactyly-long thumb syndrome	319239	Brazilian hemorrhagic fever	528	Brunzell syndrome
1277	Brachydactyly-mesomelia-intellectual disability-heart defects syndrome	1945	BRE	→528	Brunzell syndrome
1246	Brachydactyly-nystagmus-cerebellar ataxia syndrome	85284	BRESEK syndrome	47	Bruton type agammaglobulinemia
1278	Brachydactyly-preaxial hallux varus syndrome	85284	BRESHECK syndrome	528	BSCL
2956	Brachydactyly-scoliosis-carpal fusion syndrome	65682	BRIC	79304	BSEP deficiency
391646	Brachydactyly-short stature-microcephaly syndrome	99960	BRIC1	1299	BSG syndrome
3168	Brachydactyly-symphalangism syndrome	99961	BRIC2	100976	BSI
93409	Brachydactyly-syndactyly, Zhao type	99960	BRIC type 1	1980	BSPDC
93394	Brachymesophalangy II and V	99961	BRIC type 2	125	BSyn
1292	Brachymorphism-onychodysplasia-dysphalangism syndrome	99990	Brill disease	65284	BTBGD
→448242	Brachyolmia type 1, Hobaek type	99990	Brill-Zinsser disease	79241	BTD deficiency
→448242	Brachyolmia type 1, Toledo type	666	Brittle bone disease	111	BTHS
93302	Brachyolmia type 2	90354	Brittle cornea syndrome	47	BTK-deficiency
93304	Brachyolmia type 3	→33364	Brittle hair syndrome, Sabinas type	2314	Buckley syndrome
448242	Brachyolmia, Hobaek/Toledo type			131	Budd-Chiari syndrome
				36258	Buerger disease
				2285	Bull-Nixon syndrome
				312	Bullous congenital ichthyosiform erythroderma
				312	Bullous congenital ichthyosiform erythroderma of Brock
				280785	Bullous DCM
				280785	Bullous diffuse cutaneous mastocytosis
				1867	Bullous dystrophy, macular type

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312	Bullous ichthyosis			141194	CAMS1
36237	Bullous impetigo	90791	CAH due to 3-beta-hydroxysteroid dehydrogenase deficiency	53719	CAMS2
33408	Bullous lichen planus			141199	CAMS3
703	Bullous pemphigoid	90795	CAH due to 11-beta-hydroxylase deficiency	3319	CAMT
→193	Buntinx-Lormans-Martin syndrome	90793	CAH due to 17-alpha-hydroxylase deficiency	1328	Camurati-Engelmann disease
98976	Buphthalmia			3261	Canale-Smith syndrome
98976	Buphthalmos	1375	CAHMR syndrome	141	Canavan disease
98976	Buphthalmus	435988	CAID syndrome	289385	Cancer diagnosed during pregnancy
543	Burkitt lymphoma	99429	CAIS	180242	Cancer of fallopian tubes
1200	Burn-McKeown syndrome	289601	Calcification of joints and arteries	71505	Cancer-associated retinopathy
353253	Burning mouth syndrome	199260	Calcified aponeurotic fibroma	2700	Cancrum oris
800	Burton skeletal dysplasia			325004	CANDLE syndrome
800	Burton syndrome	90290	Calcinosis-Raynaud phenomenon-esophageal involvement-sclerodactyly-telangiectasia syndrome	71279	CANOMAD syndrome
352763	Buschke scleredema			2233	Cantalamessa-Baldini-Ambrosi syndrome
79501	Buschke-Fischer-Brauer syndrome	280062	Calciphylaxis	1335	Cantrell deformity
1306	Buschke-Ollendorff syndrome	280065	Calciphylaxis cutis	1335	Cantrell syndrome
99001	Butterfly-shaped pattern dystrophy			363705	Cantu craniofaciofrontodigital syndrome
99001	Butterfly-shaped pigment dystrophy	1416	Calcium pyrophosphate dihydrate crystal deposition disease	171881	Cap disease
99001	Butterfly-shaped pigmentary macular dystrophy	1408	Calderón-González-Cantu syndrome	160148	Cap inflammatory polyposis
1307	Buttiens-Fryns syndrome	228123	California disease	171881	Cap myopathy
132	Butyrylcholinesterase deficiency	83483	Californian encephalitis	160148	Cap polyposis
275864	bv-FTD	289601	CALJA	85199	CAP syndrome
1243	BVMD	85192	Calvarial doughnut lesions-bone fragility syndrome	166260	Capdepont teeth
116	BWS	→1466	CAMAK syndrome	75327	CAPE dystrophy
79306	Byler disease	3003	Camera syndrome	75327	CAPED
459353	C1 inhibitor deficiency	2163	Camero-Lituanian-Cohen syndrome		
280133	C3 deficiency	→1466	CAMFAK syndrome	188	Capillary hyperpermeability syndrome
→329931	C3 deposition glomerulonephritis without proliferation	79395	Camisa disease	188	Capillary leak syndrome
329931	C3 glomerulonephritis	83472	CAMOS syndrome	79490	Capillary lymphangioma
329918	C3 glomerulopathy	1318	Campomelia, Cumming type	79490	Capillary lymphatic malformation
401901	C9ORF72-related Huntington disease phenotype	140	Campomelic dwarfism		
401901	C9ORF72-related Huntington disease-like syndrome	140	Campomelic dysplasia	137667	Capillary malformation-arteriovenous malformation syndrome
1308	C syndrome	1319	Camptobrachydactyly	1171	CAPOS syndrome
231242	C-beta-thalassemia	295016	Camptodactyly of fingers	171839	Capra-DeMarco syndrome
97297	C-like syndrome	1327	Camptodactyly syndrome, Guadalajara type 1	464343	CAPS
401948	CA-VA deficiency	1326	Camptodactyly syndrome, Guadalajara type 2	71505	CAR syndrome
85293	Cabezas syndrome			199354	CARASIL
1309	Cacchi-Ricci disease	2848	Camptodactyly-arthropathy-coxavara-pericarditis syndrome	147	Carbamoyl-phosphate synthetase 1 deficiency
75377	CACD			147	Carbamoyl-phosphate synthetase deficiency
135	CACH syndrome	376	Camptodactyly-cleft palate-clubfoot syndrome	147	Carbamoyl-phosphate synthetase I deficiency
2848	CACP syndrome				
159	CACT deficiency	1321	Camptodactyly-fibrous tissue hyperplasia-skeletal dysplasia syndrome	79328	Carbohydrate deficient glycoprotein syndrome type 1L
56425	CAD				
448010	CAD-CDG	1323	Camptodactyly-joint contractures-facial skeletal defects syndrome	79318	Carbohydrate deficient glycoprotein syndrome type Ia
136	CADASIL				
369942	CADDS	3447	Camptodactyly-overgrowth-unusual facies syndrome	79319	Carbohydrate deficient glycoprotein syndrome type Ib
1578	CADH deficiency				
2566	CAEBV syndrome	85164	Camptodactyly-tall stature-scoliosis-hearing loss syndrome	79320	Carbohydrate deficient glycoprotein syndrome type Ic
1310	Caffey disease				
436174	CAGSSS	1325	Camptodactyly-aurinuria syndrome		
		1766	CAMRQ syndrome		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
79321	Carbohydrate deficient glycoprotein syndrome type Id	329178	Carbohydrate deficient glycoprotein syndrome type Iu	111	Cardioskeletal myopathy-neutropenia syndrome
79322	Carbohydrate deficient glycoprotein syndrome type Ie	370924	Carbohydrate deficient glycoprotein syndrome type Ix	3238	Cardiospondylocarpofacial syndrome
79323	Carbohydrate deficient glycoprotein syndrome type If	370927	Carbohydrate deficient glycoprotein syndrome type Iy	2072	Cardiovascular Gaucher disease
79324	Carbohydrate deficient glycoprotein syndrome type Ig	448010	Carbohydrate deficient glycoprotein syndrome type Iz	1358	Carey-Fineman-Ziter syndrome
				79403	Carmi syndrome
79325	Carbohydrate deficient glycoprotein syndrome type Ih	306686	Carbon monoxide-induced parkinsonism	→293843	Carnevale syndrome
79326	Carbohydrate deficient glycoprotein syndrome type Ii	2785	Carbonic anhydrase 2 deficiency	2947	Carnevale-Hernández-del Castillo syndrome
		213605	Carcinofibroma of the corpus uteri	→293843	Carnevale-Krajewska-Fischetto syndrome
397941	Carbohydrate deficient glycoprotein syndrome type II due to MAN1B1 deficiency	100093	Carcinoid syndrome	1359	Carney complex
		319308	Carcinoma associated with MITF/TFE translocation	319340	Carney complex variant
79329	Carbohydrate deficient glycoprotein syndrome type IIa	418945	Carcinoma of esophagus, salivary gland type	319340	Carney complex-trismus-pseudocamptodactyly syndrome
79330	Carbohydrate deficient glycoprotein syndrome type IIb	423781	Carcinoma of stomach, salivary gland type	97286	Carney dyad
				1359	Carney syndrome
79332	Carbohydrate deficient glycoprotein syndrome type IIc	300557	Carcinoma of the ampulla of Vater	139411	Carney triad
				213787	Carcinosarcoma of the cervix uteri
79333	Carbohydrate deficient glycoprotein syndrome type IIe	213610	Carcinosarcoma of the corpus uteri	97286	Carney-Stratakis syndrome
238459	Carbohydrate deficient glycoprotein syndrome type IIg	369891	Cardiac anomalies-developmental delay-facial dysmorphism syndrome	42	Carnitine deficiency secondary to medium-chain acyl-CoA dehydrogenase deficiency
		137628	Cardiac anomalies-heterotaxy syndrome	156	Carnitine palmitoyl transferase 1A deficiency
95428	Carbohydrate deficient glycoprotein syndrome type IIh	168796	Cardiac conduction disease-dilated cardiomyopathy-brachydactyly syndrome	228302	Carnitine palmitoyl transferase deficiency type 2, adult-onset form
263487	Carbohydrate deficient glycoprotein syndrome type III	1686	Cardiac diverticulum	228305	Carnitine palmitoyl transferase deficiency type 2, hepatocardiomyopathy form
		208600	Cardiac papillary fibroelastoma		
263501	Carbohydrate deficient glycoprotein syndrome type IIj	875	Cardiac tumor of child	228308	Carnitine palmitoyl transferase deficiency type 2, lethal systemic form
314667	Carbohydrate deficient glycoprotein syndrome type IIk	2872	Cardiocranial syndrome, Pfeiffer type		
468699	Carbohydrate deficient glycoprotein syndrome type IIl	37553	Cardiodysrhythmic potassium-sensitive periodic paralysis	228302	Carnitine palmitoyl transferase deficiency type 2, myopathic form
		1340	Cardiofaciocutaneous syndrome	228308	Carnitine palmitoyl transferase deficiency type 2, neonatal form
468684	Carbohydrate deficient glycoprotein syndrome type IIo	97292	Cardiogenic shock		
466703	Carbohydrate deficient glycoprotein syndrome type IIp	2229	Cardiogenital syndrome	228305	Carnitine palmitoyl transferase deficiency type 2, severe infantile form
		1342	Cardiomelic syndrome type 3		
86309	Carbohydrate deficient glycoprotein syndrome type Ij	500	Cardiomyopathic lentiginosis	156	Carnitine palmitoyl transferase IA deficiency
79327	Carbohydrate deficient glycoprotein syndrome type Ik	70474	Cardiomyopathy with hypotonia due to cytochrome C oxidase deficiency		
91131	Carbohydrate deficient glycoprotein syndrome type Im	70474	Cardiomyopathy with myopathy due to COX deficiency	228305	Carnitine palmitoyl transferase II deficiency, hepatocardiomyopathy form
244310	Carbohydrate deficient glycoprotein syndrome type In	1345	Cardiomyopathy-cataract-hip spine disease syndrome		
263494	Carbohydrate deficient glycoprotein syndrome type Io	91130	Cardiomyopathy-hypotonia-lactic acidosis syndrome	228308	Carnitine palmitoyl transferase II deficiency, lethal systemic form
280071	Carbohydrate deficient glycoprotein syndrome type Ip	90022	Cardiomyopathy-renal anomalies syndrome	228302	Carnitine palmitoyl transferase II deficiency, myopathic form
300536	Carbohydrate deficient glycoprotein syndrome type Ir	111	Cardioskeletal myopathy with neutropenia and abnormal mitochondria	228308	Carnitine palmitoyl transferase II deficiency, neonatal form
				228305	Carnitine palmitoyl transferase II deficiency, severe infantile form

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
157	Carnitine palmitoyltransferase deficiency type 2	→1466	Cataract-microcephaly-failure to thrive-kyphoscoliosis syndrome	468684	CCDC115-CDG
				98972	CCDF
157	Carnitine palmitoyltransferase II deficiency	1377	Cataract-microcornea syndrome	48431	CCFDN
				2008	CCGE syndrome
158	Carnitine transporter defect	2712	Cataract-microphthalmia-radiculomegaly-cardiac septal defect syndrome	99827	CCHF
158	Carnitine uptake deficiency			661	CCHS
159	Carnitine-acylcarnitine translocase deficiency	1380	Cataract-nephropathy-encephalopathy syndrome	289499	CCMCO
				319276	CCRCC
1361	Carnosinase deficiency	100990	Cataracts-motor neuropathy-short stature-skeletal anomalies syndrome	2505	CCSF
1361	Carnosinemia			457246	CCSK
53035	Caroli disease	464343	Catastrophic antiphospholipid syndrome	280779	CCV
480520	Caroli syndrome			86870	CD4+/CD56+ hematodermic neoplasm
65759	Carpenter syndrome	464343	Catastrophic APS	437552	CD16 deficiency
93973	Carpenter-Waziri syndrome	567	CATCH 22		
2767	Carpotarsal osteochondromatosis	3286	Catecholaminergic polymorphic ventricular tachycardia	238505	CD27 deficiency
64692	Carrion disease			800	Catel-Hempel syndrome
175	Cartilage-hair hypoplasia	1388	Catel-Manzke syndrome	293825	CDA due to KLF1 mutation
→175	Cartilage-hair hypoplasia-like-skeletal dysplasia without hypotrichosis syndrome	228337	Cathepsin D deficiency	98869	CDA I
				60015	Catlin marks
65282	Carvajal syndrome	85164	CATSHL syndrome	98870	CDA III
209908	CAS	1123	Caudal appendage-deafness syndrome	293825	CDA IV
56425	CAS			98869	CDA type 1
94095	Casamassima-Morton-Nance syndrome	1756	Caudal duplication	98873	CDA type 2
		3027	Caudal dysplasia	98870	CDA type 3
275517	Caspase 8 deficiency syndrome	3027	Caudal regression sequence	293825	CDA type 4
1101	Cassia Stocco dos Santos syndrome	99994	Causalgia	98869	CDA type I
160	Castleman disease	1329	CAVC	98873	CDA type II
2513	Castro Gago-Pombo-Novio syndrome	99066	CAVC type A	98870	CDA type III
		99067	CAVC type B	293825	CDA type IV
		99068	CAVC type C	85199	CDAGS syndrome
50839	Cat-scratch disease	99068	CAVC-Fallot tetralogy syndrome	293825	CDAN4
926	Catalase deficiency	99066	CAVC-left heart obstruction syndrome	247203	CDC
1373	Cataract-aberrant oral frenula-growth delay syndrome			99067	CAVC-ventricle hypoplasia syndrome
1366	Cataract-alopecia-sclerodactyly syndrome	2124	Cavernous hemangiomas of face-supraumbilical midline raphe syndrome	79319	CDG1B
1368	Cataract-ataxia-deafness syndrome			79320	CDG1C
314993	Cataract-congenital heart disease-neural tube defect syndrome	79489	Cavernous lymphangioma	79321	CDG1D
				79489	Cavernous lymphatic malformation
1383	Cataract-deafness-hypogonadism syndrome	165958	Cavitary myiasis	79323	CDG1F
162	Cataract-glaucoma syndrome	567	Cayler cardiofacial syndrome	79324	CDG1G
436174	Cataract-growth hormone deficiency-sensory neuropathy-sensorineural hearing loss-skeletal dysplasia syndrome	94122	Cayman ataxia	79325	CDG1H
		363972	CBL syndrome	79326	CDG1I
		79282	CblC defect	86309	CDG1J
		79283	CblD defect	79327	CDG1K
1375	Cataract-hypertrichosis-intellectual disability syndrome	79284	CblF defect	79328	CDG1L
		369955	CblJ defects	91131	CDG1M
1381	Cataract-intellectual disability-anal atresia-urinary defects syndrome	79284	CblF defect	244310	CDG1N
		70567	CCA	263494	CDG1O
1387	Cataract-intellectual disability-hypogonadism syndrome	115	CCA syndrome	280071	CDG1P
		2444	CCAM	324737	CDG1Q
→1466	Cataract-microcephaly-arthrogryposis-kyphosis syndrome	280832	CCAM type 1	300536	CDG1R
		280840	CCAM type 2	324422	CDG1S
		280847	CCAM type 3	319646	CDG1T
				329178	CDG1U

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
370921	CDG1W	300536	CDG syndrome type Ir	35173	CDPX2
370924	CDG1X	324422	CDG syndrome type Is	35173	CDPXD
370927	CDG1Y	319646	CDG syndrome type It	158	CDSP
448010	CDG1Z	329178	CDG syndrome type Iu	468641	CEAS
79329	CDG2A	370921	CDG syndrome type Iw	1459	CEC
79330	CDG2B	370924	CDG syndrome type Ix	2718	Cecato de Lima-Pinheiro syndrome
99843	CDG2C	370927	CDG syndrome type Iy	1515	CED
79332	CDG2D	448010	CDG syndrome type Iz	66631	CEDNIK syndrome
79333	CDG2E	79318	CDG-Ia	275517	CEDS
238459	CDG2F	79319	CDG-Ib	1459	Celiac disease-epilepsy-cerebral calcification syndrome
263508	CDG2G	79320	CDG-Ic		
95428	CDG2H	79321	CDG-Id	293208	Celiac trunk compression syndrome
263487	CDG2I	79322	CDG-Ie	93942	Celosomia
263501	CDG2J	79323	CDG-If	3258	Cenani syndactyly
314667	CDG2K	79324	CDG-Ig	3258	Cenani-Lenz syndactyly
464443	CDG2L	79325	CDG-Ih	3258	Cenani-Lenz syndrome
356961	CDG2M	79326	CDG-Ii	75377	Central areolar choroidal dystrophy
468699	CDG2N	79329	CDG-IIa	75377	Central areolar choroidal sclerosis
468684	CDG2O	79330	CDG-IIb	75327	Central areolar pigment epithelial dystrophy
466703	CDG2P	99843	CDG-IIc		
79318	CDG syndrome type Ia	79332	CDG-IId	2431	Central bilateral macrogyria
79319	CDG syndrome type Ib	79333	CDG-IIe	98972	Central cloudy corneal dystrophy of François
79320	CDG syndrome type Ic	238459	CDG-IIf	98972	Central cloudy dystrophy of François
79321	CDG syndrome type Id	263508	CDG-IIg	661	Central congenital hypoventilation syndrome
79322	CDG syndrome type Ie	95428	CDG-IIh	597	Central core disease
79323	CDG syndrome type If	263487	CDG-IIi	178029	Central diabetes insipidus
79324	CDG syndrome type Ig	263501	CDG-IIj	→98967	Central discoid corneal dystrophy
79325	CDG syndrome type Ih	314667	CDG-IIk	99832	Central hypothyroidism due to TRH receptor deficiency
79326	CDG syndrome type Ii	464443	CDG-IIl	3240	Central nervous system calcification-deafness-tubular acidosis-anemia syndrome
79329	CDG syndrome type IIa	356961	CDG-IIm		
79330	CDG syndrome type IIb	468699	CDG-IIn	73256	Central neurocytoma
99843	CDG syndrome type IIc	468684	CDG-IIo	637	Central neurofibromatosis
79332	CDG syndrome type IId	466703	CDG-IIp	295004	Central polydactyly of fingers
79333	CDG syndrome type IIe	86309	CDG-Ij	295173	Central polydactyly of fingers, bilateral
238459	CDG syndrome type IIf	79327	CDG-Ik		
263508	CDG syndrome type IIg	79328	CDG-Il	295171	Central polydactyly of fingers, unilateral
95428	CDG syndrome type IIh	91131	CDG-Ilm		
263487	CDG syndrome type IIf	244310	CDG-Iln	295010	Central polydactyly of foot
263501	CDG syndrome type IIj	263494	CDG-Ilo	295004	Central polydactyly of hand
314667	CDG syndrome type IIk	280071	CDG-Ilp	295010	Central polydactyly of toes
464443	CDG syndrome type IIl	324737	CDG-Ilq	295185	Central polydactyly of toes, bilateral
356961	CDG syndrome type IIm	300536	CDG-Ilr	295183	Central polydactyly of toes, unilateral
468699	CDG syndrome type IIn	324422	CDG-Ils		
468684	CDG syndrome type IIo	319646	CDG-Ilt	759	Central precocious puberty
466703	CDG syndrome type IIp	329178	CDG-Ilu		
86309	CDG syndrome type Ij	370921	CDG-Ilw	75327	Central retinal pigment epithelial dystrophy
79327	CDG syndrome type Ik	370924	CDG-Ilx		
79328	CDG syndrome type IIl	370927	CDG-Ily	411527	Central retinal vein occlusion
91131	CDG syndrome type IIm	448010	CDG-Ilz		
244310	CDG syndrome type IIn	2140	CDH	443079	Central serous chorioretinopathy
263494	CDG syndrome type IIo	1529	CDHS	90156	Centrifugal lipodystrophy
280071	CDG syndrome type IIp	178029	CDI	89841	Centripetal dystrophic epidermolysis bullosa
324737	CDG syndrome type IIq	1490	CDPD		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
89841	Centripetal recessive dystrophic epidermolysis bullosa	77261	Cerebral juvenile and adult form of Gaucher disease	213812	Cervical peripheral neuroectodermal cancer
89841	Centripetalis recessive dystrophic epidermolysis bullosa	221126	Cerebral proliferative glomeruloid vasculopathy	213802	Cervical rhabdomyosarcoma
319160	Centronuclear myopathy type 4	329217	Cerebral sinovenous thrombosis	268392	Cervical spina bifida aperta
1945	Centrotemporal epilepsy	447788	Cerebral visual impairment	268762	Cervical spina bifida cystica
79277	CEP	397922	Cerebro-cutaneous syndrome with iron overload	213767	Cervical squamous cell carcinoma
2398	Cephalothoracic lipodystrophy			141067	Cervicofacial enchondroma
79506	CEPT deficiency	→2995	Cerebro-oculo-facial-lymphatic syndrome	141067	Cervicofacial fibrochondroma
276183	Cerebellar ataxia with azoospermia and intellectual disability	→2995	Cerebro-oculo-facial-lymphatic syndrome	3456	Cervicooculoacoustic syndrome
94122	Cerebellar ataxia, Cayman type	1393	Cerebrocostomandibular syndrome	268397	Cervicothoracic spina bifida aperta
1171	Cerebellar ataxia-areflexia-pes cavus-optic atrophy-sensorineural hearing loss syndrome	141194	Cerebrofacial arteriovenous metameric syndrome type 1	268766	Cervicothoracic spina bifida cystica
		53719	Cerebrofacial arteriovenous metameric syndrome type 2	195	CES
1174	Cerebellar ataxia-ectodermal dysplasia syndrome	141199	Cerebrofacial arteriovenous metameric syndrome type 3	586	CF
1173	Cerebellar ataxia-hypogonadism syndrome	314679	Cerebrofacioarticular syndrome	2032	CFA
		1394	Cerebrofaciothoracic dysplasia	1340	CFC syndrome
1766	Cerebellar ataxia-intellectual disability-dysequilibrium syndrome	912	Cerebrohepatorenal syndrome	1520	CFND
		2406	Cerebromedullospinal disconnection	1520	CFNS
83472	Cerebellar ataxia-intellectual disability-optic atrophy-skin abnormalities syndrome	1458	Cerebrohepatorenal syndrome	2020	CFTDM
		1466	Cerebrooculodentofacial-skeletal syndrome	379	CGD
97249	Cerebellar atrophy with progressive microcephaly	912	Cerebrohepatorenal syndrome	2026	CGHT
		66625	Cerebrooculofacioskeletal syndrome	2388	ChAc
2246	Cerebellar hypoplasia-tapetoretinal degeneration syndrome	1396	Cerebrorenodigital syndrome	307766	CHAC syndrome
251931	Cerebellar liponeurocytoma	313838	Cerebroretinal microangiopathy with calcifications and cysts	307766	CHACS
94147	Cerebellar syndrome-pigmentary maculopathy syndrome	→247691	Cerebroretinal vasculopathy	3386	Chagas disease
		909	Cerebrotendinous xanthomatosis	436159	CHAI
1454	Cerebellar vermiform hypoplasia-oligophrenia-congenital ataxia-coloboma-hepatic fibrosis	1980	Cerebrovascular ferrocaldinosis	→1071	CHAND syndrome
		169079	Cernunnos deficiency	98979	Chandler syndrome
444072	Cerebellar-facial-dental syndrome	169079	Cernunnos XLFD	→1071	CHANDS
444072	Cerebellofaciodental syndrome	169079	Cernunnos-XLF deficiency	2235	Chang-Davidson-Carlson syndrome
2318	Cerebellooculorenal syndrome	98989	Cerulean cataract	88642	Channelopathy-associated CIP
475	Cerebelloparenchymal disorder IV	213772	Cervical adenocarcinoma	88642	Channelopathy-associated congenital insensitivity to pain
1532	Cerebellotrigeminal-dermal dysplasia syndrome	213828	Cervical adenoid basal carcinoma	3282	Chaotic atrial tachycardia
		213823	Cervical adenoid cystic carcinoma	319244	Chapare hemorrhagic fever
46724	Cerebral arteriovenous malformation	213792	Cervical adenoid cystic carcinoma	46627	Char syndrome
		99079	Cervical adenosarcoma	1964	Char-Douglas-Dungan syndrome
136	Cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy	213787	Cervical aortic arch	803	Charcot disease
		141046	Cervical carcinosarcoma	101081	Charcot-Marie-Tooth disease type 1A
199354	Cerebral autosomal recessive arteriopathy with subcortical infarcts and leukoencephalopathy	213837	Cervical germ cell cancer	101082	Charcot-Marie-Tooth disease type 1B
		2218	Cervical hypertrichosis-peripheral neuropathy syndrome	101083	Charcot-Marie-Tooth disease type 1C
66631	Cerebral dysgenesis-neuropathy-ichthyosis-palmoplantar keratoderma syndrome	213807	Cervical leiomyosarcoma	101084	Charcot-Marie-Tooth disease type 1D
		213837	Cervical malignant germ cell tumor	90658	Charcot-Marie-Tooth disease type 1E
821	Cerebral gigantism	213787	Cervical malignant Müllerian mixed tumor	101085	Charcot-Marie-Tooth disease type 1F
→1900	Cerebral gigantism, Nevo type	213812	Cervical malignant peripheral neuroectodermal tumor	98856	Charcot-Marie-Tooth disease type 2B1
2081	Cerebral gigantism-jaw cysts syndrome	213817	Cervical papillary carcinoma	101101	Charcot-Marie-Tooth disease type 2B2

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
101102	Charcot-Marie-Tooth disease type 2H	167	Chédiak-Higashi disease	101000	Childhood-onset spastic paraparesis-distal muscle wasting syndrome
		167	Chédiak-Higashi syndrome		
300319	Charcot-Marie-Tooth disease type 2P	381	Chédiak-Higashi-like syndrome	3474	CHIME syndrome
397968	Charcot-Marie-Tooth disease type 2R	167	Chédiak-Higashi-Steinbrink syndrome	2888	Chitayat-Meunier-Hodgkinson syndrome
		293603	CHEDII	3218	Chitty-Hall-Baraitser syndrome
443073	Charcot-Marie-Tooth disease type 2S	1221	Cheilitis glandularis	757	Chloride shunt syndrome
		99647	Cheirospndyloenchondromatosis	86850	Chloroma
443950	Charcot-Marie-Tooth disease type 2T	955	Cheney syndrome	180	CHM
64748	Charcot-Marie-Tooth disease type 3	812	Cherry-red spot-myoclonus syndrome	137914	Choanal atresia
99948	Charcot-Marie-Tooth disease type 4A	184	Cherubism	137920	Choanal atresia, bilateral
		3019	Cherubism-gingival fibromatosis-intellectual disability syndrome	137917	Choanal atresia, unilateral
99955	Charcot-Marie-Tooth disease type 4B1	→672	CHHS	1200	Choanal atresia-hearing loss-cardiac defects-craniofacial dysmorphism syndrome
99956	Charcot-Marie-Tooth disease type 4B2	268882	Chiari malformation type 1	70567	Cholangiocarcinoma
		1136	Chiari malformation type 2	480501	Choledochal cyst
363981	Charcot-Marie-Tooth disease type 4B3	268882	Chiari malformation type I	69663	Cholelithiasis with ABCB4 gene mutation
		1136	Chiari malformation type II	173	Cholera
99949	Charcot-Marie-Tooth disease type 4C	324625	Chikungunya		
		90280	Chilblain lupus	79303	Cholestasis with delta(4)-3-oxosteroid 5-beta-reductase deficiency
99950	Charcot-Marie-Tooth disease type 4D	139	CHILD nevus		
		139	CHILD syndrome	1414	Cholestasis-lymphedema syndrome
99951	Charcot-Marie-Tooth disease type 4E	64280	Childhood absence epilepsy	1415	Cholestasis-pigmentary retinopathy-cleft palate syndrome
		439175	Childhood AIS		
99952	Charcot-Marie-Tooth disease type 4F	209908	Childhood apraxia of speech	→2697	Cholestatic jaundice-renal tubular insufficiency syndrome
		439175	Childhood arterial ischemic stroke	75234	Cholesterol ester storage disease
99953	Charcot-Marie-Tooth disease type 4G		Childhood ataxia with diffuse central nervous system hypomyelination	79506	Cholesterol-ester transfer protein deficiency
99954	Charcot-Marie-Tooth disease type 4H	135		75234	Cholesteryl ester storage disease
139515	Charcot-Marie-Tooth disease type 4J	168782	Childhood disintegrative disorder	79344	Chondrodysplasia punctata, Sheffield type
		293955	Childhood encephalopathy due to thiamine pyrophosphokinase deficiency	79346	Chondrodysplasia punctata, tibial-metacarpal type
391351	Charcot-Marie-Tooth disease type 4K	391497	Childhood myasthenia gravis	79347	Chondrodysplasia punctata, Toriello type
90120	Charcot-Marie-Tooth disease type 6		Childhood-onset autosomal recessive myopathy with external ophthalmoplegia	263463	Chondrodysplasia with congenital joint dislocations, CHST3 type
363981	Charcot-Marie-Tooth disease with focally folded myelin	363677		280586	Chondrodysplasia with joint dislocations, gPAPP type
90658	Charcot-Marie-Tooth disease-deafness syndrome		Childhood-onset autosomal recessive slowly progressive spinocerebellar ataxia	3144	Chondrodysplasia with snail-like pelvis
		284324		50945	Chondrodysplasia, Blomstrand type
90103	Charcot-Marie-Tooth disease-deafness-intellectual disability syndrome	33402	Childhood-onset HCC	2098	Chondrodysplasia, Grebe type
		33402	Childhood-onset hepatocellular carcinoma	166272	Chondrodysplasia-dentinogenesis imperfecta-joint laxity syndrome
93114	Charcot-Marie-Tooth disease-nephropathy syndrome	247667	Childhood-onset hypophosphatasia	1422	Chondrodysplasia-disorder of sex development syndrome
64751	Charcot-Marie-Tooth disease-pyramidal features syndrome	171439	Childhood-onset nemaline myopathy	1422	Chondrodysplasia-pseudohermaphroditism syndrome
138	CHARGE association		Childhood-onset phosphoethanolaminuria	35173	Chondrodystrophia calcificans congenita
138	CHARGE syndrome	247667			
921	CHARGE-like syndrome		Childhood-onset progressive contractures-limb-girdle weakness-muscle dystrophy syndrome		
1496	Charlevoix disease	466921			
1406	Charlie M syndrome		Childhood-onset Rathburn disease		
168577	CHC type 2	247667			
98975	CHED1				
293603	CHED2				
98975	CHEDI				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
289	Chondroectodermal dysplasia	2137	Chronic autoimmune hepatitis	439202	Chronic obstetric brachial plexus injury
319195	Chondroectodermal dysplasia with night blindness	133	Chronic berylliosis	439202	Chronic obstetric brachial plexus palsy
		133	Chronic beryllium disease		
404507	Chondromyxoid fibroma	133	Chronic beryllium lung disease	95426	Chronic pain requiring intraspinal analgesia
55880	Chondrosarcoma	56425	Chronic cold agglutinin disease	330064	Chronic photosensitivity dermatitis
444077	CHOPS syndrome	79078	Chronic dacryoadenitis and sialadenitis	91359	Chronic pneumonitis of infancy
251674	Chordoid glioma				
178	Chordoma	103907	Chronic diarrhea due to glucoamylase deficiency	324964	Chronic recurrent multifocal osteomyelitis
2388	Chorea-acanthocytosis				
2388	Choreoacanthocytosis	314373	Chronic diarrhea due to guanylate cyclase 2C overactivity	77297	Chronic recurrent multifocal osteomyelitis-congenital dyserythropoietic anemia-neutrophilic dermatosis syndrome
209905	Choreoathetosis-hypothyroidism-neonatal respiratory distress syndrome	397606	Chronic diarrhea with hereditary sensory and autonomic neuropathy		
252015	Choriocarcinoma of the central nervous system	397606	Chronic diarrhea with HSAN	217566	Chronic respiratory distress with surfactant metabolism deficiency
251899	Choroid plexus carcinoma	1670	Chronic diarrhea with villous atrophy		
2807	Choroid plexus papilloma	2566	Chronic EBV infection syndrome	71279	Chronic sensory ataxic neuropathy with anti-dialosyl IgM antibodies
1433	Choroidal atrophy-alopecia syndrome	279891	Chronic endophthalmitis	379	Chronic septic granulomatosis
		468641	Chronic enteropathy associated with SLCO2A1 gene	83418	Chronic spinal muscular atrophy
39044	Choroidal melanoma	168940	Chronic eosinophilic leukemia	70591	Chronic thromboembolic pulmonary hypertension
180	Choroideremia				
1435	Choroideremia-deafness-obesity syndrome	2902	Chronic eosinophilic pneumonia	97353	Chronic traumatic encephalopathy
1434	Choroideremia-hypopituitarism syndrome	2566	Chronic Epstein-Barr virus infection syndrome	37748	Chronic urticaria with gammopathy
		99921	Chronic graft versus host disease	37748	Chronic urticaria with macroglobulinemia
94087	CHP	521	Chronic granulocytic leukemia	263463	CHST3-related skeletal dysplasia
181	Christ-Siemens-Touraine syndrome	379	Chronic granulomatous disease	2953	CHST14-related EDS
1436	Christian syndrome	396	Chronic hiccup	2953	CHST14-related Ehlers-Danlos syndrome
85278	Christianson syndrome	1451	Chronic infantile neurological cutaneous articular syndrome	93971	Chudley-Lowry syndrome
1808	Christianson-Fourie syndrome			93971	Chudley-Lowry-Hoar syndrome
98879	Christmas disease	83418	Chronic infantile spinal muscular atrophy	314597	Chudley-McCullough syndrome
182	Chromoblastomycosis			3068	Chudley-Rozdilsky syndrome
182	Chromomycosis	2932	Chronic inflammatory demyelinating polyneuropathy	183	Churg-Strauss syndrome
319303	Chromophobe renal cell adenocarcinoma	2932	Chronic inflammatory demyelinating polyradiculoneuropathy	238557	Chuvash erythrocytosis
319303	Chromophobe renal cell carcinoma			238557	Chuvash polycythemia
1450	Chromosome 8-derived supernumerary ring/marker	294422	Chronic intestinal failure	71	Chylomicron retention disease
3380	Chromosome 18 duplication	2978	Chronic intestinal pseudoobstruction	1160	Chylous ascites
330064	Chronic actinic dermatitis			46486	Cicatrical pemphigoid
314928	Chronic adult hydrocephalus	284448	Chronic lymphocytic inflammation with pontine perivascular enhancement responsive to steroids	217390	CID due to DOCK8 deficiency
99871	Chronic and localized Langerhans cell histiocytosis			317473	CID due to IKAROS deficiency
137817	Chronic arachnoiditis	1334	Chronic mucocutaneous candidiasis	445018	CID due to LRBA deficiency
		99873	Chronic multifocal Langerhans cell histiocytosis	317476	CID due to MAGT1 deficiency
71279	Chronic ataxic neuropathy-ophthalmoplegia-IgM paraprotein-cold agglutinins-disialosyl antibodies syndrome	521	Chronic myelogenous leukemia	317428	CID due to ORAI1 deficiency
		521	Chronic myeloid leukemia	231154	CID due to partial RAG1 deficiency
		98823	Chronic myelomonocytic leukemia	443811	CID due to PGM3 deficiency
435988	Chronic atrial and intestinal dysrhythmia syndrome	86830	Chronic myeloproliferative disease, unclassifiable	157949	CID due to RAG 1/2 deficiency
435988	Chronic atrial dysrhythmia-intestinal motility disorder	77261	Chronic neuronopathic Gaucher disease	317430	CID due to STIM1 deficiency
325004	Chronic atypical neutrophilic dermatosis-lipodystrophy-elevated temperature syndrome	86829	Chronic neutrophilic leukemia	314689	CID due to STK4 deficiency
				231154	CID with expansion of gamma delta T cells
				436252	CID-MIA/early-onset IBD
				435651	CIDEC-related familial partial lipodystrophy

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
435651	CIDEC-related FPLD	315311	Classic congenital adrenal hyperplasia due to 21-hydroxylase deficiency, simple virilizing form	485350	CLCN4-related X-linked intellectual disability syndrome
2932	CIDP			398971	Clear cell adenocarcinoma of ovary
79394	CIE			404511	Clear cell papillary renal cell carcinoma
294422	CIF	325524	Classic congenital lipoid adrenal hyperplasia due to STAR deficiency	319276	Clear cell renal carcinoma
1223	Ciliary dysentery	93930	Classic exstrophy of the bladder	319276	Clear cell renal cell adenocarcinoma
2114	Cilliers-Beighton syndrome	79239	Classic galactosemia	319276	Clear cell renal cell carcinoma
1451	CINCA syndrome	98962	Classic GCD	319276	Clear cell renal cell carcinoma
391397	CIP with hyperhidrosis and gastrointestinal dysfunction	289857	Classic glycine encephalopathy	457246	Clear cell sarcoma of kidney
642	CIP-anhidrosis syndrome	98962	Classic granular corneal dystrophy	97338	Clear cell sarcoma of the tendons and aponeuroses
478664	CIP-hypohidrosis syndrome	391	Classic Hodgkin disease		
2978	CIPO	391	Classic Hodgkin lymphoma	101023	Cleft hard palate
69744	Circumscribed palmoplantar hypokeratosis	98846	Classic Hodgkin lymphoma, lymphocyte-depleted type	141291	Cleft lip and alveolus
309854	Cirrhosis-dystonia-polycythemia-hypermanganesemia syndrome	98845	Classic Hodgkin lymphoma, lymphocyte-rich type	199306	Cleft lip and palate
				199306	Cleft lip-alveolus-palate syndrome
57777	Cirrhotic cardiomyopathy	98844	Classic Hodgkin lymphoma, mixed cellularity type	1995	Cleft lip-retinopathy syndrome
157820	CISS			199306	Cleft lip/palate
247525	Citrullinemia type 1	98843	Classic Hodgkin lymphoma, nodular sclerosis type	888	Cleft lip/palate with mucous cysts of lower lip
247585	Citrullinemia type 2			2319	Cleft lip/palate-abnormal thumbs-microcephaly syndrome
247525	Citrullinemia type I	394	Classic homocystinuria	2003	Cleft lip/palate-deafness-sacral lipoma syndrome
247585	Citrullinemia type II	475	Classic Joubert syndrome		
251383	CK syndrome	313	Classic lamellar ichthyosis	3253	Cleft lip/palate-ectodermal dysplasia syndrome
90790	CLAH	98964	Classic lattice corneal dystrophy		
97249	CLAM	268145	Classic maple syrup urine disease	→1896	Cleft lip/palate-ectrodactyly syndrome
168984	CLAPO syndrome	158796	Classic mast cell leukemia		
188	Clarkson disease	251867	Classic medulloblastoma	2328	Cleft lip/palate-facial, eye, heart and intestinal anomalies syndrome
466026	Class I G6PD deficiency	324604	Classic MmD	2001	Cleft lip/palate-intestinal malrotation-cardiopathy syndrome
466026	Class I glucose-6-phosphate dehydrogenase deficiency	268145	Classic MSUD		
90794	Classic 21-OHD CAH	324604	Classic multiminicore disease	3253	Cleft lip/palate-syndactyly-pili torti syndrome
315306	Classic 21-OHD CAH, salt wasting form	324604	Classic multiminicore myopathy		
		2584	Classic mycosis fungoides	95465	Cleft mitral valve
315311	Classic 21-OHD CAH, simple virilizing form	329977	Classic neuroendocrine tumor of appendix	141242	Cleft nose
85138	Classic Addison's disease	216866	Classic pantothenate kinase-associated neurodegeneration	2008	Cleft palate-cardiac defect-genital anomalies-ectrodactyly syndrome
329977	Classic appendiceal neuroendocrine tumor	163898	Classic paraneoplastic limbic encephalitis	921	Cleft palate-coloboma-deafness syndrome
329977	Classic appendix neuroendocrine tumor	163898	Classic paraneoplastic limbic encephalitis, with or without intracellular antigens	2013	Cleft palate-large ears-small head syndrome
93605	Classic Bartter syndrome	93258	Classic Pfeiffer syndrome	2016	Cleft palate-lateral synechia syndrome
268145	Classic BCKD deficiency	79254	Classic phenylketonuria	2015	Cleft palate-short stature-vertebral anomalies syndrome
268145	Classic branched-chain 2-ketoacid dehydrogenase deficiency	79254	Classic PKU		
268145	Classic branched-chain ketoaciduria	280219	Classic PMD	2010	Cleft palate-stapes fixation-oligodontia syndrome
247525	Classic citrullinemia	240071	Classic progressive supranuclear palsy syndrome		
247546	Classic citrullinemia type 1	240071	Classic PSP syndrome	99772	Cleft soft palate
247546	Classic citrullinemia type I	773	Classic Refsum disease	99772	Cleft velum
325524	Classic CLAH	18	Classic RTA	99772	Cleft velum palatinum
90794	Classic congenital adrenal hyperplasia due to 21-hydroxylase deficiency	443192	Classic SPS	3429	Cleft-limb-heart malformation syndrome syndrome
		443192	Classic stiff person syndrome		
315306	Classic congenital adrenal hyperplasia due to 21-hydroxylase deficiency, salt wasting form	3467	Classic xanthinuria	1997	Clefting-ectropion-conical teeth syndrome
		2272	Clayton Smith-Donnai syndrome	1452	Cleidocranial dysostosis

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
1452	Cleidocranial dysplasia	521	CML	99952	CMT4F
3472	Cleidocranial dysplasia-micrognathia-absent thumbs syndrome	98823	CMML	99953	CMT4G
		252202	CMMR-D syndrome	99954	CMT4H
		99763	CMO I	139515	CMT4J
1453	Cleidorhizomelic syndrome	99763	CMO II	391351	CMT4K
284448	CLIPPERS	238459	CMP-sialic acid transporter deficiency	101078	CMT4X
228329	CLN1 disease			99014	CMT5X
228349	CLN2 disease			90120	CMT6
228346	CLN3 disease	71	CMRD	352675	CMT6X
228340	CLN4A disease	590	CMS	1556	CMTC
228343	CLN4B disease	101081	CMT1A	100043	CMTDIA
228360	CLN5 disease	101082	CMT1B	100044	CMTDIB
228363	CLN6 disease	101083	CMT1C	100045	CMTDIC
228366	CLN7 disease	101084	CMT1D	100046	CMTDID
228354	CLN8 disease	90658	CMT1E	93114	CMTDIE
1947	CLN8 disease, Northern epilepsy variant	101085	CMT1F	352670	CMTDIF
		101075	CMT1X	101075	CMTX1
228357	CLN9 disease	324611	CMT2 due to KIF5A mutation	101076	CMTX2
228337	CLN10 disease	435819	CMT2 due to TFG mutation	101077	CMTX3
314629	CLN11 disease	435387	CMT2 due to VCP mutation	101078	CMTX4
314632	CLN12 disease	401964	CMT2 with giant axons	99014	CMTX5
352709	CLN13 disease	99946	CMT2A1	352675	CMTX6
93929	Cloacal exstrophy	99947	CMT2A2	468635	CMUSE
314950	Clonal hypereosinophilic syndrome	99936	CMT2B	137698	CMV disease in patients with impaired cell mediated immunity deemed at risk
221083	Clonic hemifacial spasm	99937	CMT2C		
268366	Closed iniencephaly	99938	CMT2D		
189	Clouston syndrome	99939	CMT2E	319160	CNM4
100978	Cloverleaf skull-asphyxiating thoracic dysplasia syndrome	99940	CMT2F	306686	CO-induced parkinsonism
		99941	CMT2G	1454	COACH syndrome
93274	Cloverleaf skull-micromelic bone dysplasia syndrome	101102	CMT2H	1456	Coarctation of the abdominal aorta
		99942	CMT2I	397725	COASY protein-associated neurodegeneration
93267	Cloverleaf skull-multiple congenital anomalies syndrome	99943	CMT2J		
		99944	CMT2K	313838	Coats plus syndrome
140944	CLOVES syndrome	99945	CMT2L	79282	Cobalamin C defect
411493	CLP1-related pontocerebellar hypoplasia	228179	CMT2M	79283	Cobalamin D defect
		228174	CMT2N	79284	Cobalamin F defect
3253	CLPED1	284232	CMT2O	369955	Cobalamin J defect
192	CLS	300319	CMT2P	53721	Cobb syndrome
85136	CLWM	329258	CMT2Q	352682	Cobblestone lissencephaly without muscular or eye involvement
137667	CM-AVM	397968	CMT2R		
289504	CMAMMA	443073	CMT2S	352682	Cobblestone lissencephaly without muscular or ocular involvement
1334	CMC	443950	CMT2T		
258	CMD1A	397735	CMT2U	1911	Cocaine embryofetopathy
98893	CMD1B	447964	CMT2V	90068	Cocaine intoxication
→370953	CMD1C	466775	CMT2X	228123	Coccidioides infection
370959	CMD with cerebellar involvement	466768	CMT2Z	228123	Coccidioidomycosis
370968	CMD with intellectual disability	101077	CMT3X	3233	Cochleasaccular degeneration-cataract syndrome
329178	CMD with intellectual disability and severe epilepsy	99948	CMT4A		
370980	CMD without intellectual disability	99955	CMT4B1	191	Cockayne syndrome
370959	CMD-CRB	99956	CMT4B2	90321	Cockayne syndrome type 1
370968	CMD-MR	363981	CMT4B3	90322	Cockayne syndrome type 2
370980	CMD-no MR	99949	CMT4C	90324	Cockayne syndrome type 3
371007	CMDH	99950	CMT4D	90321	Cockayne syndrome type I
		99951	CMT4E		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
90322	Cockayne syndrome type II	98947	Coloboma of optic papilla	79283	Combined defect in adenosylcobalamin and methylcobalamin synthesis, type cbID
90324	Cockayne syndrome type III	155884	Coloboma of superior eyelid		
1458	CODAS syndrome	3474	Coloboma-congenital heart disease-ichthyosiform dermatosis-intellectual disability-ear anomalies syndrome	79284	Combined defect in adenosylcobalamin and methylcobalamin synthesis, type cbIF
192	Coffin-Lowry syndrome				
1465	Coffin-Siris syndrome				
1466	COFS syndrome				
263508	COG1-CDG	138	Coloboma-heart defects-atresia choanae-retardation of growth and development-genitourinary problems-ear abnormalities syndrome	369955	Combined defect in adenosylcobalamin and methylcobalamin synthesis, type cbIJ
435934	COG2-CDG				
435934	COG2-related congenital disorder of glycosylation				
263501	COG4-CDG				
263487	COG5-CDG				
464443	COG6-CDG	468672	Colobomatous macrophthalmia-microcornea syndrome	369962	Combined defect in adenosylcobalamin and methylcobalamin synthesis, type cbIX
79333	COG7-CDG	98938	Colobomatous microphthalmia		
95428	COG8-CDG	363741	Colobomatous microphthalmia-obesity-hypogenitalism-intellectual disability syndrome	35909	Combined deficiency of factor V and factor VIII
1467	Cogan syndrome				
98980	Cogan-Reese syndrome	424099	Colobomatous microphthalmia-rhizomelic dysplasia syndrome	99732	Combined deficiency of sulfite oxidase, xanthine dehydrogenase and aldehyde oxidase
444077	Cognitive impairment-coarse facies-heart defects-obesity-pulmonary involvement-short stature-skeletal dysplasia syndrome	435930	Colobomatous optic disc-macular atrophy-chorioretinopathy syndrome		
193	Cohen syndrome	→138	Colobomatous-microphthalmia-heart disease-hearing loss syndrome	308386	Combined deficiency of sulfite oxidase, xanthine dehydrogenase and aldehyde oxidase type A
2969	Cohen-Hayden syndrome	1198	Colonic atresia	308393	Combined deficiency of sulfite oxidase, xanthine dehydrogenase and aldehyde oxidase type B
79144	COIF	100080	Colonic NET		
79144	COIF syndrome	16	Color blindness, blue monochrome monochromatic type	308400	Combined deficiency of sulfite oxidase, xanthine dehydrogenase and aldehyde oxidase type C
36383	COL4A1-related brain small vessel disease with hemorrhage	83595	Colorado tick encephalitis		
36383	COL4A1-related familial vascular leukoencephalopathy	83595	Colorado tick fever	440727	Combined hamartoma of the retina and retinal pigment epithelium
36383	COL4A1-related retinal arteriolar tortuosity-infantile hemiparesis-autosomal dominant leukoencephalopathy syndrome	83595	Colorado tick-borne disease		
		733	Colorectal adenomatous polyposis	440727	Combined hamartoma of the retina and RPE
31824	Colchicine poisoning	261584	Colorectal adenomatous polyposis due to monosomy 5q22.2	221078	Combined hyperactive dysfunction syndrome of the cranial nerves
56425	Cold agglutinin disease	90793	Combined 17-hydroxylase/17,20-lyase deficiency	169082	Combined immunodeficiency due to CD3gamma deficiency
56425	Cold agglutinin syndrome	445062	Combined cerebellar and peripheral ataxia-hearing loss-diabetes mellitus syndrome	169090	Combined immunodeficiency due to CRAC channel dysfunction
157820	Cold-induced sweating syndrome				
324561	Cole disease	370114	Combined cervical dystonia	217390	Combined immunodeficiency due to dedicator of cytokinesis 8 protein deficiency
2050	Cole-Carpenter syndrome				
84087	Collagen type III glomerulopathy	356978	Combined D-2-hydroxyglutaric acidemia and L-2-hydroxyglutaric acidemia	217390	Combined immunodeficiency due to DOCK8 deficiency
84087	Collagenofibrotic glomerulopathy				
247203	Collecting duct carcinoma	356978	Combined D-2-hydroxyglutaric aciduria and L-2-hydroxyglutaric aciduria	317473	Combined immunodeficiency due to IKAROS deficiency
2412	Collins-Pope syndrome				
98942	Coloboma of choroid and retina	26	Combined defect in adenosylcobalamin and methylcobalamin synthesis	445018	Combined immunodeficiency due to LRBA deficiency
98943	Coloboma of eye lens				
98946	Coloboma of eyelid	79282	Combined defect in adenosylcobalamin and methylcobalamin synthesis, type cbIC	317476	Combined immunodeficiency due to MAGT1 deficiency
155889	Coloboma of inferior eyelid				
98944	Coloboma of iris	397964		317428	Combined immunodeficiency due to ORAI1 deficiency
98945	Coloboma of macula				
1471	Coloboma of macula-brachydactyly type B syndrome	1475	Coloboma of optic nerve with renal disease	431149	Combined immunodeficiency due to OX40 deficiency
98947	Coloboma of optic disc				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
231154	Combined immunodeficiency due to partial RAG1 deficiency	314637	Combined oxidative phosphorylation defect type 10	280133	Complement component 3 deficiency
443811	Combined immunodeficiency due to PGM3 deficiency	324535	Combined oxidative phosphorylation defect type 11	99429	Complete androgen insensitivity syndrome
157949	Combined immunodeficiency due to RAG 1/2 deficiency	314051	Combined oxidative phosphorylation defect type 12	99429	Complete androgen resistance syndrome
317430	Combined immunodeficiency due to STIM1 deficiency	319514	Combined oxidative phosphorylation defect type 13	1329	Complete atrioventricular canal
314689	Combined immunodeficiency due to STK4 deficiency	319519	Combined oxidative phosphorylation defect type 14	99066	Complete atrioventricular canal type A
911	Combined immunodeficiency due to ZAP70 deficiency	319524	Combined oxidative phosphorylation defect type 15	99067	Complete atrioventricular canal type B
431149	Combined immunodeficiency with childhood-onset Kaposi sarcoma	352563	Combined oxidative phosphorylation defect type 16	99068	Complete atrioventricular canal type C
231154	Combined immunodeficiency with expansion of gamma delta T cells	369913	Combined oxidative phosphorylation defect type 17	99066	Complete atrioventricular canal-left heart obstruction syndrome
221139	Combined immunodeficiency with faciooculoskeletal anomalies	420728	Combined oxidative phosphorylation defect type 20	99068	Complete atrioventricular canal-tetralogy of Fallot syndrome
39041	Combined immunodeficiency with hypereosinophilia	420733	Combined oxidative phosphorylation defect type 21	99067	Complete atrioventricular canal-ventricle hypoplasia syndrome
431149	Combined immunodeficiency with impaired immunity to HHV-8	444013	Combined oxidative phosphorylation defect type 23	1329	Complete atrioventricular septal defect
431149	Combined immunodeficiency with impaired immunity to human herpes virus 8	444458	Combined oxidative phosphorylation defect type 24	98949	Complete cryptophthalmia
157949	Combined immunodeficiency with skin granulomas	447954	Combined oxidative phosphorylation defect type 25	289916	Complete deficiency of methylmalonyl-CoA mutase
228423	Combined immunodeficiency with susceptibility to mycobacterial, viral and fungal infections	477684	Combined oxidative phosphorylation defect type 26	633	Complete growth hormone insensitivity
436252	Combined immunodeficiency-enteropathy spectrum	477774	Combined oxidative phosphorylation defect type 27	254688	Complete hydatidiform mole
169079	Combined immunodeficiency-microcephaly-growth retardation-sensitivity to ionizing radiation syndrome	466784	Combined oxidative phosphorylation defect type 28	79293	Complete LCAT deficiency
1979	Combined insulin, insulin-like growth factor 1 (IGF1) and epidermal growth factor (EGF) deficiency	478029	Combined oxidative phosphorylation defect type 29	29	Complete mevalonate kinase deficiency
289504	Combined malonic and methylmalonic acidemia	478042	Combined oxidative phosphorylation defect type 30	254688	Complete molar pregnancy
289504	Combined malonic and methylmalonic aciduria	309111	Combined pancreatic lipase-colipase deficiency	49382	Complete or incomplete color blindness
254920	Combined oxidative phosphorylation defect type 2	95494	Combined pituitary hormone deficiencies, genetic forms	180126	Complete septate uterus
254925	Combined oxidative phosphorylation defect type 4	139406	Combined prosaposin deficiency	101063	Complete situs inversus
137908	Combined oxidative phosphorylation defect type 5	300564	Combined pulmonary fibrosis-emphysema syndrome	101063	Complete situs inversus viscerum
254930	Combined oxidative phosphorylation defect type 7	166286	Comedo nevus of the palm	180074	Complete unilateral aplasia of the Müllerian ducts
319504	Combined oxidative phosphorylation defect type 8	141276	Commissural facial cleft	180074	Complete unilateral Müllerian aplasia
319509	Combined oxidative phosphorylation defect type 9	141061	Commissural lip fistula	457378	Complex lethal osteochondrodysplasia
		3384	Common aorticopulmonary trunk	457378	Complex lethal osteochondrodysplasia, Symoens-Barnes-Gistelink type
		3384	Common arterial trunk	83452	Complex regional pain syndrome
		1329	Common atrioventricular canal	99995	Complex regional pain syndrome type 1
		→288	Common hereditary elliptocytosis	99994	Complex regional pain syndrome type 2
		620	Common mesentery	306644	Complication after organ transplantation
		1572	Common variable immunodeficiency	268316	Complication in hemodialysis
		280821	Communicating congenital bronchopulmonary-foregut malformation	458758	Composite hemangioendothelioma
				168966	Composite Hodgkin and non-Hodgkin lymphoma

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
168966	Composite lymphoma	295091	Congenital absence of thigh and lower leg with foot present, bilateral	3319	Congenital amegakaryocytic thrombocytopenia
634	Comèl-Netherton syndrome			3319	Congenital amegakaryocytic thrombocytopenic purpura
228165	Concentric demyelination	295089	Congenital absence of thigh and lower leg with foot present, unilateral	86816	Congenital analbuminemia
383	Conductive deafness with stapes fixation			217399	Congenital analgesia with hyperhidrosis
3216	Conductive deafness-malformed external ear syndrome	93322	Congenital absence of tibia	453510	Congenital analgesia with severe intellectual disability
3236	Conductive deafness-ptosis-skeletal anomalies syndrome	435623	Congenital absence of toes		
3216	Conductive hearing loss-malformed external ear syndrome	295118	Congenital absence of toes, bilateral	95507	Congenital anomaly of hepatic vein
1871	Cone dystrophy	295116	Congenital absence of toes, unilateral	91489	Congenital anterior megalophthalmia
209932	Cone dystrophy with supernormal rod electroretinogram	2879	Congenital absence of ulna and fibula		
209932	Cone dystrophy with supernormal rod ERG	294975	Congenital absence of upper arm and forearm with hand present	95449	Congenital aortic valve insufficiency
209932	Cone dystrophy with supernormal rod response	295087	Congenital absence of upper arm and forearm with hand present, bilateral	3093	Congenital aortic valve stenosis
209932	Cone dystrophy with supernormal scotopic electroretinogram			2037	Congenital aortopulmonary artery fistula
1872	Cone rod dystrophy	295085	Congenital absence of upper arm and forearm with hand present, unilateral	2037	Congenital aortopulmonary septal defect
1873	Cone rod dystrophy-amelogenesis imperfecta syndrome	247775	Congenital absence of uterus and vagina	2037	Congenital aortopulmonary window
221142	Confetti-like macular atrophy			93322	Congenital aplasia and dysplasia of the tibia with intact fibula
440233	Congenital abducens nerve palsy	96269	Congenital absence of vagina	353334	Congenital arteriovenous anastomoses of the retina
294979	Congenital absence of both forearm and hand	294990	Congenital absence/hypoplasia of fingers excluding thumb	353334	Congenital arteriovenous communication of the retina
295095	Congenital absence of both forearm and hand, bilateral	295114	Congenital absence/hypoplasia of fingers excluding thumb, bilateral	1195	Congenital atransferrinemia
295093	Congenital absence of both forearm and hand, unilateral	973	Congenital absence/hypoplasia of fingers excluding thumb, unilateral	60041	Congenital atrioventricular block
294981	Congenital absence of both lower leg and foot	294988	Congenital absence/hypoplasia of thumb	162526	Congenital auditory ossicle malformation without external ear abnormality
295099	Congenital absence of both lower leg and foot, bilateral	295112	Congenital absence/hypoplasia of thumb, bilateral	1216	Congenital benign spinal muscular atrophy with contractures
295097	Congenital absence of both lower leg and foot, unilateral	295110	Congenital absence/hypoplasia of thumb, unilateral	48	Congenital bilateral absence of vas deferens
289465	Congenital absence of fingerprints	324353	Congenital achiasma	48	Congenital bilateral agenesis of vas deferens
294986	Congenital absence of foot	93583	Congenital ADAMTS-13 deficiency	48	Congenital bilateral aplasia of vas deferens
295107	Congenital absence of foot, bilateral	90791	Congenital adrenal hyperplasia due to 3-beta-hydroxysteroid dehydrogenase deficiency	93177	Congenital bilateral megacalycosis
295105	Congenital absence of foot, unilateral	90795	Congenital adrenal hyperplasia due to 11-beta-hydroxylase deficiency	79301	Congenital bile acid synthesis defect type 1
294983	Congenital absence of hand	90793	Congenital adrenal hyperplasia due to 17-alpha-hydroxylase deficiency	79303	Congenital bile acid synthesis defect type 2
295103	Congenital absence of hand, bilateral	95699	Congenital adrenal hyperplasia due to cytochrome P450 oxidoreductase deficiency	79302	Congenital bile acid synthesis defect type 3
295101	Congenital absence of hand, unilateral	95699	Congenital adrenal hyperplasia due to cytochrome POR deficiency	79095	Congenital bile acid synthesis defect type 4
86815	Congenital absence of lacrimal puncta and salivary glands	95701	Congenital adrenal hypoplasia of maternal cause	2292	Congenital bowing of long bones
217399	Congenital absence of pain with hyperhidrosis	33355	Congenital aleukocytosis	71278	Congenital brain dysgenesis due to glutamine synthetase deficiency
453510	Congenital absence of pain with severe intellectual disability	79	Congenital alpha2-antiplasmin deficiency	2040	Congenital bronchobiliary fistula
294977	Congenital absence of thigh and lower leg with foot present	210122	Congenital alveolar capillary dysplasia	3161	Congenital bronchopulmonary sequestration
				289499	Congenital cataract microcornea with corneal opacity

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
300313	Congenital cataract-deafness-severe developmental delay syndrome	280840	Congenital cystic adenomatoid malformation of the lung type 2	91131	Congenital disorder of glycosylation type 1m
300313	Congenital cataract-hearing loss-severe developmental delay syndrome	280847	Congenital cystic adenomatoid malformation of the lung type 3	244310	Congenital disorder of glycosylation type 1n
		280854	Congenital cystic adenomatoid malformation of the lung type 4	263494	Congenital disorder of glycosylation type 1o
1369	Congenital cataract-hypertrophic cardiomyopathy-mitochondrial myopathy syndrome	2444	Congenital cystic adenomatous malformation of the lung	280071	Congenital disorder of glycosylation type 1p
1376	Congenital cataract-ichthyosis syndrome	280827	Congenital cystic adenomatous malformation of the lung type 0	324737	Congenital disorder of glycosylation type 1q
464738	Congenital cataract-microcephaly-nevus flammeus simplex-severe intellectual disability syndrome	280832	Congenital cystic adenomatous malformation of the lung type 1	300536	Congenital disorder of glycosylation type 1r
		280840	Congenital cystic adenomatous malformation of the lung type 2	324422	Congenital disorder of glycosylation type 1s
330054	Congenital cataract-progressive muscular hypotonia-deafness-developmental delay syndrome	280847	Congenital cystic adenomatous malformation of the lung type 3	329178	Congenital disorder of glycosylation type 1u
		280854	Congenital cystic adenomatous malformation of the lung type 4	370921	Congenital disorder of glycosylation type 1w
330054	Congenital cataract-progressive muscular hypotonia-hearing loss-developmental delay syndrome	480501	Congenital cystic dilatation of the biliary tract	370924	Congenital disorder of glycosylation type 1x
48431	Congenital cataracts-facial dysmorphism-neuropathy syndrome	2444	Congenital cystic disease of the lung	370927	Congenital disorder of glycosylation type 1y
661	Congenital central alveolar hypoventilation syndrome	280832	Congenital cystic disease of the lung type 1	448010	Congenital disorder of glycosylation type 1z
99803	Congenital central alveolar hypoventilation-Hirschsprung disease syndrome	280840	Congenital cystic disease of the lung type 2	397941	Congenital disorder of glycosylation type 2 due to MAN1B1 deficiency
2345	Congenital cervical vertebral fusion	280847	Congenital cystic disease of the lung type 3	79329	Congenital disorder of glycosylation type 2a
53689	Congenital chloride diarrhea	168612	Congenital deficiency in alpha-fetoprotein	79330	Congenital disorder of glycosylation type 2b
329242	Congenital chronic diarrhea with exudative enteropathy	2140	Congenital diaphragmatic hernia	79332	Congenital disorder of glycosylation type 2d
329242	Congenital chronic diarrhea with protein-losing enteropathy	3474	Congenital disorder of glycosylation due to PIGL deficiency	79333	Congenital disorder of glycosylation type 2e
264688	Congenital chylothorax	79318	Congenital disorder of glycosylation type 1a	238459	Congenital disorder of glycosylation type 2f
2505	Congenital circumferential skin folds	79319	Congenital disorder of glycosylation type 1b	263508	Congenital disorder of glycosylation type 2g
91413	Congenital Claude-Bernard-Horner syndrome	79320	Congenital disorder of glycosylation type 1c	95428	Congenital disorder of glycosylation type 2h
440221	Congenital CNIII lesion	79321	Congenital disorder of glycosylation type 1d	263487	Congenital disorder of glycosylation type 2i
98686	Congenital CNIV palsy	79322	Congenital disorder of glycosylation type 1e	263501	Congenital disorder of glycosylation type 2j
440233	Congenital CNVI palsy	79323	Congenital disorder of glycosylation type 1f	314667	Congenital disorder of glycosylation type 2k
269505	Congenital communicating hydrocephalus	79324	Congenital disorder of glycosylation type 1g	464443	Congenital disorder of glycosylation type 2l
99129	Congenital complete agenesis of pericardium	79325	Congenital disorder of glycosylation type 1h	356961	Congenital disorder of glycosylation type 2m
115	Congenital contractural arachnodactyly	79326	Congenital disorder of glycosylation type 1i	468699	Congenital disorder of glycosylation type 2n
178382	Congenital convex foot	86309	Congenital disorder of glycosylation type 1j	468684	Congenital disorder of glycosylation type 2o
178382	Congenital convex pes valgus	79327	Congenital disorder of glycosylation type 1k	466703	Congenital disorder of glycosylation type 2p
53691	Congenital cornea plana	79328	Congenital disorder of glycosylation type 1L		
95491	Congenital coronary artery aneurysm				
2444	Congenital cystic adenomatoid malformation of the lung				
280827	Congenital cystic adenomatoid malformation of the lung type 0				
280832	Congenital cystic adenomatoid malformation of the lung type 1				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
79318	Congenital disorder of glycosylation type Ia	244310	Congenital disorder of glycosylation type In	231573	Congenital erosive and vesicular dermatosis
79319	Congenital disorder of glycosylation type Ib	263494	Congenital disorder of glycosylation type Io	90042	Congenital erythrocytosis due to erythropoietin receptor mutation
79320	Congenital disorder of glycosylation type Ic	280071	Congenital disorder of glycosylation type Ip	369992	Congenital erythroderma-hypotrichosis-recurrent infections-multiple food allergies syndrome
79321	Congenital disorder of glycosylation type Id	324737	Congenital disorder of glycosylation type Iq		79277
79322	Congenital disorder of glycosylation type Ie	300536	Congenital disorder of glycosylation type Ir	91358	Congenital esophageal diverticulum
79323	Congenital disorder of glycosylation type If	324422	Congenital disorder of glycosylation type Is	215	Congenital essential nyctalopia
				91	Congenital estrogen deficiency
79324	Congenital disorder of glycosylation type Ig	329178	Congenital disorder of glycosylation type Iu	280811	Congenital extrapulmonary sequestration
79325	Congenital disorder of glycosylation type Ih	370921	Congenital disorder of glycosylation type Iw	99176	Congenital eyelid retraction
79326	Congenital disorder of glycosylation type Ii	370924	Congenital disorder of glycosylation type Ix	570	Congenital facial diplegia
				325	Congenital factor II deficiency
397941	Congenital disorder of glycosylation type II due to MAN1B1 deficiency	370927	Congenital disorder of glycosylation type Iy	326	Congenital factor V deficiency
				327	Congenital factor VII deficiency
79329	Congenital disorder of glycosylation type IIa	293825	Congenital dyserythropoietic anemia due to KLF1 mutation	328	Congenital factor X deficiency
				329	Congenital factor XI deficiency
79330	Congenital disorder of glycosylation type IIb	98869	Congenital dyserythropoietic anemia type 1	330	Congenital factor XII deficiency
				331	Congenital factor XIII deficiency
79332	Congenital disorder of glycosylation type IIc	98873	Congenital dyserythropoietic anemia type 2	2020	Congenital fiber-type disproportion myopathy
79333	Congenital disorder of glycosylation type IIe	98870	Congenital dyserythropoietic anemia type 3	335	Congenital fibrinogen deficiency
238459	Congenital disorder of glycosylation type IIg	293825	Congenital dyserythropoietic anemia type 4	45358	Congenital fibrosis of extraocular muscles
				90045	Congenital folate malabsorption
263508	Congenital disorder of glycosylation type IIh	98869	Congenital dyserythropoietic anemia type I	98686	Congenital fourth cranial nerve palsy
95428	Congenital disorder of glycosylation type IIi	98873	Congenital dyserythropoietic anemia type II	2345	Congenital fused cervical segments
263487	Congenital disorder of glycosylation type IIj	98870	Congenital dyserythropoietic anemia type III	476406	Congenital generalized hypercontractile muscle stiffness syndrome
263501	Congenital disorder of glycosylation type IIk	293825	Congenital dyserythropoietic anemia type IV	2026	Congenital generalized hypertrichosis terminalis
314667	Congenital disorder of glycosylation type IIl	67044	Congenital dyserythropoietic anemia with thrombocytopenia	1023	Congenital generalized hypertrichosis, Ambras type
464443	Congenital disorder of glycosylation type IIl	70596	Congenital EBV infection	79495	Congenital generalized hypertrichosis, Macias-Flores type
		91491	Congenital ectropion uveae	295232	Congenital genu flexum
356961	Congenital disorder of glycosylation type IIm	295032	Congenital elbow dislocation	295229	Congenital genu recurvatum
468699	Congenital disorder of glycosylation type IIo	295227	Congenital elbow dislocation, bilateral	157826	Congenital gingival cell tumor
		295225	Congenital elbow dislocation, unilateral	98976	Congenital glaucoma
468684	Congenital disorder of glycosylation type IIo	103910	Congenital enterocyte heparan sulfate deficiency	157826	Congenital granular cell tumor
466703	Congenital disorder of glycosylation type IIp	168601	Congenital enterokinase deficiency	330	Congenital Hageman factor deficiency
86309	Congenital disorder of glycosylation type IIq	168601	Congenital enteropathy due to enteropeptidase deficiency	60041	Congenital heart block
79327	Congenital disorder of glycosylation type IIr	292	Congenital enterovirus infection	1355	Congenital heart defect-round face-developmental delay syndrome
		70596	Congenital Epstein-Barr virus infection	139	Congenital hemidysplasia with ichthyosiform nevus and limbs defects
91131	Congenital disorder of glycosylation type IIr	157826	Congenital epulis		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
98975	Congenital hereditary endothelial dystrophy type 1	231662	Congenital IGHD type IA	70472	Congenital lactic acidosis, Saguenay-Lac-Saint-Jean type
		231671	Congenital IGHD type IB		
293603	Congenital hereditary endothelial dystrophy type 2	231679	Congenital IGHD type II	313	Congenital lamellar ichthyosis
		231692	Congenital IGHD type III		
98975	Congenital hereditary endothelial dystrophy type I	306504	Congenital ILNEB syndrome	99872	Congenital Langerhans cell histiocytosis
293603	Congenital hereditary endothelial dystrophy type II	217399	Congenital indifference to pain with hyperhidrosis	141124	Congenital laryngeal cyst
				137932	Congenital laryngeal palsy
306530	Congenital hereditary facial palsy with variable deafness	64752	Congenital insensitivity to pain and thermal analgesia	2374	Congenital laryngeal web
				2373	Congenital laryngomalacia
306530	Congenital hereditary facial palsy with variable hearing loss	217399	Congenital insensitivity to pain with hyperhidrosis	1954	Congenital lethal erythroderma
				210163	Congenital lethal myopathy, Compton-North type
306530	Congenital hereditary facial paralysis with variable deafness	391397	Congenital insensitivity to pain with hyperhidrosis and gastrointestinal dysfunction	93937	Congenital limb amputation
				90790	Congenital lipoid adrenal hyperplasia due to STAR deficiency
306530	Congenital hereditary facial paralysis-variable deafness syndrome	453510	Congenital insensitivity to pain with preserved temperature sensation		
		453510	Congenital insensitivity to pain with severe intellectual disability	140944	Congenital lipomatous overgrowth-vascular malformation-epidermal nevi-skeletal anomaly syndrome
306530	Congenital hereditary facial paralysis-variable hearing loss syndrome				
		453510	Congenital insensitivity to pain with severe non-progressive cognitive delay	140944	Congenital lipomatous overgrowth-vascular malformation-epidermal nevi-spinal anomaly syndrome
101068	Congenital hereditary stromal dystrophy			1928	Congenital lobar emphysema
293	Congenital herpes simplex virus infection	642	Congenital insensitivity to pain-anhidrosis syndrome	1928	Congenital lobar hyperinflation
				768	Congenital long QT syndrome
483	Congenital high-molecular-weight kininogen deficiency	478664	Congenital insensitivity to pain-hypohidrosis syndrome		
91413	Congenital Horner syndrome			93323	Congenital longitudinal deficiency of the fibula
2185	Congenital hydrocephalus	306504	Congenital interstitial lung disease-nephrotic syndrome-epidermolysis bullosa syndrome		
2190	Congenital hydronephrosis			93321	Congenital longitudinal deficiency of the radius
478	Congenital hypogonadotropic hypogonadism with anosmia	388	Congenital intestinal aganglionosis		
		280802	Congenital intrapulmonary sequestration	93322	Congenital longitudinal deficiency of the tibia
124	Congenital hypoplastic anemia, Blackfan-Diamond type	1229	Congenital intrauterine infection-like syndrome	93320	Congenital longitudinal deficiency of the ulna
→672	Congenital hypothalamic hamartoma syndrome	332	Congenital intrinsic factor deficiency	2430	Congenital macroglossia
		199296	Congenital isolated ACTH deficiency	95430	Congenital major airway collapse
226313	Congenital hypothyroidism due to maternal intake of antithyroid drugs	631	Congenital isolated GH deficiency		
		231662	Congenital isolated GH deficiency type IA	83620	Congenital malabsorptive diarrhea due to paucity of enteroendocrine cells
95715	Congenital hypothyroidism due to transplacental passage of maternal TSH-binding inhibitory antibodies	231671	Congenital isolated GH deficiency type IB	93109	Congenital megacalycosis
		231679	Congenital isolated GH deficiency type II	280671	Congenital megaconial myopathy
1195	Congenital hypotransferrinemia				
→113	Congenital hypotrichosis-milia syndrome	231679	Congenital isolated GH deficiency type II	69063	Congenital membranous nephropathy due to maternal anti-neutral endopeptidase alloimmunization
		231692	Congenital isolated GH deficiency type III		
88621	Congenital ichthyosis type 4			2665	Congenital mesoblastic nephroma
352333	Congenital ichthyosis-intellectual disability-spastic quadriplegia syndrome	631	Congenital isolated growth hormone deficiency	621	Congenital methemoglobinemia
		231662	Congenital isolated growth hormone deficiency type IA		
352333	Congenital ichthyosis-intellectual disability-spastic tetraplegia syndrome	231671	Congenital isolated growth hormone deficiency type IB	391376	Congenital microcephaly-severe encephalopathy-progressive cerebral atrophy syndrome
		231679	Congenital isolated growth hormone deficiency type II	566	Congenital microcoria
2271	Congenital ichthyosis-microcephalus-quadruplegia syndrome			199293	Congenital microgastria
		231692	Congenital isolated growth hormone deficiency type III	2290	Congenital microvillous atrophy
2271	Congenital ichthyosis-microcephalus-tetraplegia syndrome			2290	Congenital microvillus atrophy
		295034	Congenital knee dislocation	566	Congenital miosis
631	Congenital IGHD	53690	Congenital lactase deficiency	99057	Congenital mitral stenosis

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98905	Congenital multicore myopathy with external ophthalmoplegia	831	Congenital narrowing of cervical spinal canal	295234	Congenital patella dislocation, unilateral
258	Congenital muscular dystrophy due to laminin alpha2 deficiency	162521	Congenital nasal pyriform aperture stenosis with holoprosencephaly	99072	Congenital patent ductus arteriosus aneurysm
157973	Congenital muscular dystrophy due to LMNA mutation	168486	Congenital NCL	332	Congenital pernicious anemia
		306504	Congenital NEP syndrome	626	Congenital pigmented nevus
280671	Congenital muscular dystrophy due to phosphatidylcholine biosynthesis defect	443988	Congenital nephrosis-cerebral ventriculomegaly syndrome	465	Congenital plasminogen activator inhibitor type 1 deficiency
258	Congenital muscular dystrophy type 1A	839	Congenital nephrotic syndrome, Finnish type	2907	Congenital poikiloderma with bullae, Weary type
98893	Congenital muscular dystrophy type 1B	306504	Congenital nephrotic syndrome-epidermolysis bullosa-pulmonary disease syndrome	90042	Congenital polycythemia due to erythropoietin receptor mutation
→370953	Congenital muscular dystrophy type 1C			480531	Congenital portosystemic shunt
→370953	Congenital muscular dystrophy type 1D	306504	Congenital nephrotic syndrome-interstitial lung disease-epidermolysis bullosa syndrome	480531	Congenital portosystemic venous fistula
		168486	Congenital neuronal ceroid lipofuscinosis	124	Congenital PRCA
370959	Congenital muscular dystrophy with cerebellar involvement			749	Congenital prekallikrein deficiency
371007	Congenital muscular dystrophy with hyperlaxity	369852	Congenital neutropenia-bone marrow fibrosis-nephromegaly syndrome	83461	Congenital primary aphakia
34520	Congenital muscular dystrophy with integrin alpha-7 deficiency			79452	Congenital primary lymphedema
370968	Congenital muscular dystrophy with intellectual disability	369852	Congenital neutropenia-myelofibrosis-nephromegaly syndrome	617	Congenital primary megaloureter
				617	Congenital primary megaureter
329178	Congenital muscular dystrophy with intellectual disability and severe epilepsy	79394	Congenital non-bullous ichthyosiform erythroderma	238654	Congenital primary megaureter, nonrefluxing and unobstructed form
34520	Congenital muscular dystrophy with ITGA7 deficiency	269510	Congenital non-communicating hydrocephalus	238646	Congenital primary megaureter, obstructed form
280671	Congenital muscular dystrophy with mitochondrial structural abnormalities	269505	Congenital non-obstructive hydrocephalus	238650	Congenital primary megaureter, refluxing form
370980	Congenital muscular dystrophy without intellectual disability	1216	Congenital nonprogressive spinal muscular atrophy	327	Congenital proconvertin deficiency
272	Congenital muscular dystrophy, Fukuyama type	208513	Congenital nonprogressive spinocerebellar ataxia	66630	Congenital pseudoarthrosis of clavicle
75840	Congenital muscular dystrophy, Ullrich type	269510	Congenital obstructive hydrocephalus	295020	Congenital pseudoarthrosis of the femur
1875	Congenital muscular dystrophy-infantile cataract-hypogonadism syndrome	440221	Congenital oculomotor nerve palsy	295022	Congenital pseudoarthrosis of the fibula
590	Congenital myasthenic syndrome	79144	Congenital onychodysplasia	157808	Congenital pseudoarthrosis of the limbs
353327	Congenital myasthenic syndromes with glycosylation defect	79144	Congenital onychodysplasia of the index fingers	295024	Congenital pseudoarthrosis of the radius
98904	Congenital myopathy with excess of thin filaments	157713	Congenital or early infantile CACH syndrome	295018	Congenital pseudoarthrosis of the tibia
319160	Congenital myopathy with internal nuclei and atypical cores	2772	Congenital osteogenesis imperfecta-microcephaly-cataracts syndrome	295026	Congenital pseudoarthrosis of the ulna
424107	Congenital myopathy with myasthenic-like onset	465	Congenital PAI-1 deficiency	91411	Congenital ptosis
199329	Congenital myopathy, Paradas type	2805	Congenital pancreatic agenesis	2444	Congenital pulmonary airway malformation
168572	Congenital myopathy-cleft palate-malignant hyperthermia syndrome	313906	Congenital pancreatic cyst	280827	Congenital pulmonary airway malformation type 0
289380	Congenital myosclerosis, Löwenthal type	139414	Congenital panfollicular nevus	280832	Congenital pulmonary airway malformation type 1
		264675	Congenital PAP	280832	Congenital pulmonary airway malformation type 1
		99130	Congenital partial agenesis of pericardium	280840	Congenital pulmonary airway malformation type 2
		99124	Congenital partial pulmonary venous return anomaly	280840	Congenital pulmonary airway malformation type 2
		295036	Congenital patella dislocation	280847	Congenital pulmonary airway malformation type 3
		295237	Congenital patella dislocation, bilateral	280854	Congenital pulmonary airway malformation type 4

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264675	Congenital pulmonary alveolar proteinosis	101068	Congenital stromal corneal dystrophy	93583	Congenital thrombotic thrombocytopenic purpura
2414	Congenital pulmonary lymphangiectasia	328	Congenital Stuart factor deficiency	99125	Congenital total pulmonary venous return anomaly
		141121	Congenital subglottic stenosis		
3161	Congenital pulmonary sequestration	35122	Congenital sucrose-isomaltase deficiency	858	Congenital toxoplasmosis
				141127	Congenital tracheal stenosis
3189	Congenital pulmonary valve stenosis	306446	Congenital sucrose-isomaltase deficiency with minimal starch tolerance	3347	Congenital tracheobronchomegaly
3188	Congenital pulmonary veins atresia or stenosis			95430	Congenital tracheomalacia
		185	Congenital pulmonary venolobar syndrome	95459	Congenital tricuspid stenosis
124	Congenital pure red cell aplasia			306474	Congenital sucrose-isomaltase deficiency with starch and lactose intolerance
		295032	Congenital radial head dislocation	210576	
3269	Congenital radioulnar synostosis	306436	Congenital sucrose-isomaltase deficiency with starch intolerance	88629	Congenital tritanopia
97598	Congenital renal artery stenosis			98686	Congenital trochlear nerve palsy
97598	Congenital renovascular hypoplasia	306462	Congenital sucrose-isomaltase deficiency without starch intolerance	93583	Congenital TTP
281190	Congenital reticular ichthyosiform erythroderma			141099	Congenital tubular nose
		92050	Congenital tufting enteropathy		
353334	Congenital retinal arteriovenous anastomoses	306486	Congenital sucrose-isomaltase deficiency without sucrose intolerance	99060	Congenital unguarded mitral orifice
353334	Congenital retinal arteriovenous communication			95457	Congenital unguarded tricuspid orifice
		91495	Congenital retinal detachment	35122	Congenital sucrose-isomaltose malabsorption
190	Congenital retinal telangiectasia	306446	Congenital sucrose-isomaltose malabsorption with minimal starch tolerance	2258	Congenital unilateral pulmonary hypoplasia
178382	Congenital rocker-bottom foot			1864	Congenital valvular dysplasia
290	Congenital rubella syndrome	306474	Congenital sucrose-isomaltose malabsorption with starch and lactose intolerance	291	Congenital varicella syndrome
974	Congenital scalp defects with distal limb anomalies			2291	Congenital velopharyngeal incompetence
		974	Congenital scalp defects with distal limb reduction anomalies	178382	Congenital vertical talus
2301	Congenital short bowel syndrome	306436	Congenital sucrose-isomaltose malabsorption with starch intolerance	295203	Congenital vertical talus, bilateral
1987	Congenital short femur			295201	Congenital vertical talus, unilateral
295030	Congenital shoulder dislocation	306462	Congenital sucrose-isomaltose malabsorption without starch intolerance	137932	Congenital vocal cord paralysis
93400	Congenital sialidosis type 2			216694	Congenitally corrected transposition of the great arteries
260305	Congenital sideroblastic anemia	306446	Congenital sucrose intolerance with minimal starch tolerance	216694	Congenitally corrected transposition of the great vessels
369861	Congenital sideroblastic anemia-B-cell immunodeficiency-periodic fever-developmental delay syndrome			2391	Congenitally short costocoracoid ligament
		306474	Congenital sucrose intolerance with starch and lactose intolerance	860	Congenitally uncorrected transposition of the great arteries
263435	Congenital smooth muscle hamartoma	306436	Congenital sucrose intolerance with starch intolerance	216729	Congenitally uncorrected transposition of the great arteries with cardiac malformation
103908	Congenital sodium diarrhea	306462	Congenital sucrose intolerance without starch intolerance		
94068	Congenital spondyloepiphyseal dysplasia	306486	Congenital sucrose-isomaltose malabsorption without sucrose intolerance	99042	Congenitally uncorrected transposition of the great arteries with coarctation
215	Congenital stationary night blindness			3465	Congenital suprabulbar paresis
		99059	Congenital supra-valvular mitral ring		
75382	Congenital stationary night blindness, Oguchi type	98948	Congenital symblepharon	216729	Congenitally uncorrected transposition of the great vessels with cardiac malformation
		99856	Congenital syringomyelia		
99122	Congenital stenosis of the inferior caval vein	2039	Congenital systemic arteriovenous fistula	99042	Congenitally uncorrected transposition of the great vessels with coarctation
99122	Congenital stenosis of the inferior vena cava	210576	Congenital temporomandibular joint ankylosis		
99122	Congenital stenosis of the IVC	440221	Congenital third cranial nerve palsy	99827	Congo fever
3197	Congenital stiff man syndrome			99827	Congo hemorrhagic fever

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97566	Congo red-negative amyloidosis-like glomerulopathy	180118	Cordiform uterus	275543	Corpus callosum hypoplasia-retardation-adducted thumbs-spasticity-hydrocephalus syndrome
		366	Cori disease		
97231	Conjunctivitis lignosa	366	Cori-Forbes disease	2318	CORS
369929	Conn adenoma with seizures and neurological abnormalities	1051	Corneal anesthesia-deafness-intellectual disability syndrome	1389	Cortical blindness-intellectual disability-polydactyly syndrome
280210	Connatal PMD	1661	Corneal dystrophy epithelial-short stature syndrome	300570	Cortical dysgenesis with pontocerebellar hypoplasia due to TUBB3 mutation
300284	Connective tissue disorder due to LH3 deficiency	98962	Corneal dystrophy Groenouw type I		
300284	Connective tissue disorder due to lysyl hydroxylase-3 deficiency	98969	Corneal dystrophy Groenouw type II	268994	Cortical dysplasia, Taylor type
→2909	Connective tissue dysplasia, Spellacy type	98961	Corneal dystrophy of Bowman layer type 1	163681	Cortical dysplasia-focal epilepsy syndrome
420794	Cono-spondylar dysplasia	98960	Corneal dystrophy of Bowman layer type 2	3152	Cortical hyperostosis-syndactyly syndrome
140969	Conorenal syndrome	98961	Corneal dystrophy of Bowman layer type I	447788	Cortical visual impairment
567	Conotruncal anomaly face syndrome	98960	Corneal dystrophy of Bowman layer type II	454887	Corticobasal syndrome
35173	Conradi-Hünemann-Happle syndrome	1490	Corneal dystrophy with progressive deafness	199247	Corticosteroid-binding globulin deficiency
319651	Constitutional megaloblastic anemia with severe neurologic disease	1490	Corneal dystrophy-perceptive deafness syndrome	54251	Corticosteroid-sensitive aseptic abscess syndrome
252202	Constitutional mismatch repair deficiency syndrome	352662	Corneal intraepithelial dyskeratosis-palmoplantar hyperkeratosis-laryngeal dyskeratosis syndrome	99763	Corticosterone methyloxidase deficiency type I
295000	Constriction rings syndrome			96253	Corticotroph pituitary adenoma
1303	Constrictive bronchiolitis	3177	Corneal-cerebellar syndrome	141163	Cosack syndrome
369942	Contiguous ABCD1 DXS1357E deletion syndrome	199	Cornelia de Lange syndrome	67047	Costeff optic atrophy syndrome
84142	Continuous muscle fiber activity syndrome	96095	Cornelia de Lange-like syndrome	67047	Costeff syndrome
		3194	Corneo-dermato-osseous syndrome	3071	Costello syndrome
725	Continuous spikes and waves during sleep	2041	Coronaro-cardiac fistula	1507	Costovertebral segmentation defect-mesomelia syndrome
		2041	Coronary arterial fistulas	1914	Coumarin embryopathy
725	Continuous spikes and waves during slow-wave sleep	2041	Coronary arterial malformations	93333	Cousin syndrome
436003	Contractures-developmental delay-Pierre Robin syndrome	99085	Coronary artery intramyocardial course	1507	COVEDEM syndrome
		99118	Coronary sinus atresia	101078	Cowchock syndrome
1484	Contractures-ectodermal dysplasia-cleft lip/palate syndrome	99117	Coronary sinus stenosis	201	Cowden disease
		50	Corpus callosum agenesis of with chorioretinal abnormality	201	Cowden syndrome
314002	Contractures-webbed neck-micrognathia-hypoplastic nipples syndrome	2508	Corpus callosum agenesis-abnormal genitalia syndrome	99932	Cow's milk hypersensitivity
1487	Cooks syndrome	3338	Corpus callosum agenesis-blepharophimosis-Robin sequence syndrome	70472	COX deficiency, French-Canadian type
231214	Cooley anemia			781	Coxiellosis
1488	Cooper-Jabs syndrome	1492	Corpus callosum agenesis-double urinary collecting system syndrome	1508	Coxoauricular syndrome
1302	COP	52055	Corpus callosum agenesis-intellectual disability-coloboma-micrognathia syndrome	1509	Coxopodopatellar syndrome
444092	COPA defect			254920	COXPD2
397725	CoPAN	459074	Corpus callosum agenesis-macrocephaly-hypertelorism syndrome	254925	COXPD4
2062	Copenhagen syndrome			137908	COXPD5
98984	Coppock-like cataract	1496	Corpus callosum agenesis-neuronopathy syndrome	254930	COXPD7
457185	COQ4-related neonatal encephalomyopathy			319504	COXPD8
99098	Cor triatriatum dexter	1553	Corpus callosum agenesis-polysyndactyly syndrome	319509	COXPD9
99098	Cor triatriatum dextrum			314637	COXPD10
99099	Cor triatriatum sinister	→3157	Corpus callosum dysgenesis-hypopituitarism syndrome	324535	COXPD11
99099	Cor triatriatum sinistrum			314051	COXPD12
98990	Coralliform cataract			319514	COXPD13
				319519	COXPD14
				319524	COXPD15
				352563	COXPD16
				369913	COXPD17

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
420728	COXPD20	2115	Cranio-facio-digito-genital syndrome	1527	Craniosynostosis, Philadelphia type
420733	COXPD21			1541	Craniosynostosis, Warman type
444013	COXPD23	1525	Cranio-osteoarthropathy	1532	Craniosynostosis-alopecia-brain defect syndrome
444458	COXPD24	2053	Craniocarpotarsal dysplasia		
447954	COXPD25	2053	Craniocarpotarsal dystrophy	85199	Craniosynostosis-anal anomalies-porokeratosis syndrome
477684	COXPD26	7	Craniocerebellocardiac dysplasia		
477774	COXPD27	1513	Craniodiaphyseal dysplasia	1530	Craniosynostosis-cataract syndrome
466784	COXPD28	1514	Craniodigital syndrome-intellectual disability syndrome	2872	Craniosynostosis-congenital heart disease-intellectual disability syndrome
478029	COXPD29				
478042	COXPD30	1515	Cranioectodermal dysplasia	1538	Craniosynostosis-Dandy-Walker malformation-hydrocephalus syndrome
2444	CPAM	2099	Craniofacial and osseous defects-intellectual disability syndrome		
280827	CPAM type 0	85168	Craniofacial conodysplasia	1535	Craniosynostosis-dysmorphism-brachydactyly syndrome
280832	CPAM type 1				
280840	CPAM type 2	1681	Craniofacial duplication	1533	Craniosynostosis-fibular aplasia syndrome
280847	CPAM type 3	1777	Craniofacial dysmorphism-coloboma-corpor callosum agenesis syndrome		
280854	CPAM type 4				
475	CPD IV	→1394	Craniofacial dysmorphism-skeletal anomalies-intellectual disability syndrome	171839	Craniosynostosis-hydrocephalus-Arnold-Chiari malformation type I-radioulnar synostosis syndrome
300564	CPFE				
91359	CPI	1798	Craniofacial dysostosis-diaphyseal hyperplasia syndrome	52054	Craniosynostosis-intracranial calcifications syndrome
2016	CPLS syndrome				
2807	CPP	2095	Craniofacial dysostosis-genital, dental, cardiac anomalies syndrome	1540	Craniosynostosis-midfacial hypoplasia-foot abnormalities syndrome
759	CPP				
147	CPS1 deficiency	314555	Craniofacial dysplasia-osteopenia syndrome	1528	Craniotelencephalic dysplasia
147	CPS1D				
156	CPT1A deficiency	459061	Craniofacial dysplasia-short stature-ectodermal anomalies-intellectual disability syndrome	2095	Cranofacial dysostosis-hypertrichosis-hypoplasia of labia majora syndrome
157	CPT2				
228302	CPT2, adult-onset form	1516	Craniofacial dyssynostosis	75373	CRAPB
228305	CPT2, hepatocardiomyocardial form			1529	Craniofacial-deafness-hand syndrome
228308	CPT2, lethal systemic form	293843	Craniofacial-ulnar-renal syndrome		
228302	CPT2, myopathic form			363705	Craniofaciofrontodigital syndrome
228308	CPT2, neonatal form	1520	Craniofrontonasal dysplasia		
228305	CPT2, severe infantile form			228390	Craniofrontonasal dysplasia with alopecia and hypogonadism
157	CPTII	1519	Craniofrontonasal dysplasia, Teebi type		
228302	CPTII, adult-onset form			90290	Craniofrontonasal dysplasia-Poland anomaly syndrome
228305	CPTII, hepatocardiomyocardial form	281190	Craniofrontonasal syndrome		
228308	CPTII, lethal systemic form			205	Craniofrontonasal syndrome
228302	CPTII, myopathic form	79234	Craniofrontonasal syndrome		
228308	CPTII, neonatal form			50814	Craniofrontonasal syndrome
228305	CPTII, severe infantile form	85184	Craniofrontonasal syndrome		
3286	CPVT			1522	Craniofrontonasal syndrome
35173	CPXD	1524	Craniofrontonasal syndrome		
2081	Cramer-Niederellmann syndrome			54595	Craniofrontonasal syndrome
202	Crandall syndrome	63260	Craniofrontonasal syndrome		
1512	Crane-Heise syndrome			157832	Craniofrontonasal syndrome
97339	Cranial dural arteriovenous fistula	284149	Craniofrontonasal syndrome		
97339	Cranial dural arteriovenous malformations			1541	Craniofrontonasal syndrome
268820	Cranial meningocele	2145	Craniofrontonasal syndrome		
98919	Cranial variant of GBS			1541	Craniofrontonasal syndrome
98919	Cranial variant of Guillain-Barré syndrome	1541	Craniofrontonasal syndrome		
420485	Cranio-cervical dystonia with laryngeal and upper-limb involvement			2145	Craniofrontonasal syndrome

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2935	Crossed polydactyly	→3157	Culler-Jones syndrome	228285	Cutis laxa acquisita
2935	Crossed polysyndactyly	3207	Curatolo-Cilio-Pessagno syndrome		Cutis laxa with severe pulmonary, gastrointestinal and urinary anomalies
439881	Croupous bronchitis	98960	Curly fiber corneal dystrophy	221145	
207	Crouzon craniofacial dysostosis		Curly hair-acral keratoderma-carries syndrome	2962	Cutis laxa-corneal clouding-intellectual disability syndrome
207	Crouzon disease	307766		171719	Cutis laxa-Marfanoid syndrome
93262	Crouzon syndrome-acanthosis nigricans syndrome	→1071	Curly hair-ankyloblepharon-nail dysplasia syndrome	1556	Cutis marmorata telangiectatica congenita
93262	Crouzon-dermoskeletal syndrome	1525	Currarino disease		Cutis verticis gyrata-intellectual disability syndrome
2905	Crow-Fukase syndrome	1525	Currarino idiopathic osteoarthropathy	→357225	
290	CRS		Currarino syndrome		Cutis verticis gyrata-retinitis pigmentosa-neurosensory deafness syndrome
→247691	CRV	1552	Currarino triad	→357225	
411527	CRVO	1552	Current pressure-sensitive neuropathy	→357225	Cutis verticis gyrata-retinitis pigmentosa-neurosensory hearing loss syndrome
91139	Cryoglobulinemia type 1	640	Curry-Hall syndrome		Cutis verticis gyrata-retinitis pigmentosa-sensorineural deafness syndrome
91138	Cryoglobulinemic vasculitis	952	Curry-Jones syndrome	→357225	
1546	Cryptococcosis	1553	Cushing disease		Cutis verticis gyrata-retinitis pigmentosa-sensorineural hearing loss syndrome
2032	Cryptogenic fibrosing alveolitis	96253	Cushing syndrome due to ectopic ACTH secretion	→357225	
163708	Cryptogenic late-onset epileptic spasms	99889	Cushing syndrome due to macronodular adrenal hyperplasia		Cutis verticis gyrata-thyroid aplasia-intellectual disability syndrome
468635	Cryptogenic multifocal ulcerous stenosing enteritis	189427	Cutaneomeningospinal angiomatosis	→357225	Cutler-Bass-Romshe syndrome
1302	Cryptogenic organizing pneumonia	53721	Cutaneous and mucosal venous malformation	→357225	CVID
1547	Cryptomicrotia-brachydactyly syndrome	2451	Cutaneous collagenous vasculopathy	466670	Cyanide poisoning
1547	Cryptomicrotia-brachydactyly-excess fingertip arch syndrome	280779	Cutaneous hemangioma with muscle or bone atrophy	306692	Cyanide-induced parkinsonism
2052	Cryptophthalmos-syndactyly syndrome	329324	Cutaneous hypersensitivity vasculitis	2686	Cyclic neutropenia
1548	Cryptorchidism-arachnodactyly-intellectual disability syndrome	889	Cutaneous infectious botulism	228379	Cyclosporine-induced folliculodystrophy
1549	Cryptosporidiosis	178475	Cutaneous larva migrans	210	Cyclosporiasis
357329	Cryptosporidiosis-chronic cholangitis-liver disease syndrome	423717	Cutaneous leukocytoclastic angiitis	79493	CYLD cutaneous syndrome
98967	Crystalline stromal dystrophy	889	Cutaneous leukocytoclastic vasculitis	171886	Cylindrical spirals myopathy
101068	CSCD	79455	Cutaneous local mastocytoma	90795	CYP11B1 deficiency
443079	CSCR	79490	Cutaneous lymphangioma circumscriptum	2674	Cyprus facial-neuromusculoskeletal syndrome
35122	CSID	79455	Cutaneous mastocytoma	212	Cystathionase deficiency
306446	CSID with minimal starch tolerance	90395	Cutaneous mucinosis of infancy	212	Cystathione gamma-lyase deficiency syndrome
306474	CSID with starch and lactose intolerance	79140	Cutaneous neuroendocrine carcinoma	394	Cystathionine beta-synthase deficiency
306436	CSID with starch intolerance	439729	Cutaneous PAN	212	Cystathioninuria
306462	CSID without starch intolerance	439729	Cutaneous periarteritis nodosa	100008	Cystatin amyloidosis
306486	CSID without sucrose intolerance	2881	Cutaneous photosensitivity-lethal colitis syndrome	400	Cystic echinococcosis
1465	CSS	439729	Cutaneous polyarteritis nodosa	586	Cystic fibrosis
100008	CST3-related amyloidosis	451607	Cutaneous pseudolymphoma	2575	Cystic fibrosis-gastritis-megaloblastic anemia syndrome
329217	CSVT	889	Cutaneous small vessel vasculitis	2111	Cystic hamartoma of lung and kidney
725	CSWS	178475	Cutaneous toxin-mediated botulism	79486	Cystic hygroma
725	CSWSS syndrome	464321	Cutaneovisceral angiomatosis-thrombocytopenia syndrome	85136	Cystic leukoencephalopathy without megalencephaly
70591	CTEPH	1555	Cutis gyrata-acanthosis nigricans-craniosynostosis syndrome	229	Cystic medial necrosis of aorta
436159	CTLA-4 haploinsufficiency with autoimmune infiltration disease				
247525	CTLN1				
247585	CTLN2				
909	CTX				
158	CUD				

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1560	Cysticercosis	329457	DA5D	56304	De la Chapelle dysplasia
213	Cystinosis	1495	Da Silva syndrome	393	De la Chapelle syndrome
214	Cystinuria	251515	DA10	3157	De Morsier syndrome
93612	Cystinuria type A	458768	Dabska tumor	→910	De Sanctis-Cacchione syndrome
93613	Cystinuria type B	141083	Dacryocele	1570	De Smet-Fabry-Fryns syndrome
214	Cystinuria-lysinuria syndrome	1562	Dacryocystitis-osteopoikilosis syndrome	33355	De Vaal disease
75381	Cystoid macular dystrophy			71277	De Vivo disease
472	Cystoisosporiasis	141083	Dacryocystocele	3214	Deaf blind hypopigmentation syndrome, Yemenite type
180261	Cystosarcoma phylloide	2186	Daentl-Townsend-Siegel syndrome		
180261	Cystosarcoma phylloide	1563	Dahlberg syndrome	90024	Deafness with labyrinthine aplasia, microtia, and microdontia
70472	Cytochrome C oxidase deficiency, French-Canadian type	1563	Dahlberg-Borer-Newcomer syndrome		
70472	Cytochrome oxidase deficiency, Saguenay-Lac-Saint-Jean type	2181	Daish-Hardman-Lamont syndrome	2663	Deafness-cataracts-skeletal anomalies syndrome
95702	Cytomegalic congenital adrenal hypoplasia	275523	DALD	3241	Deafness-craniofacial syndrome
		1183	Dancing eye syndrome	52368	Deafness-dystonia-optic neuropathy syndrome
	1183	Dancing eye-dancing feet syndrome			
137698	Cytomegalovirus disease in patients with impaired cell mediated immunity deemed at risk	→42775	Dandy-Walker malformation-facial hemangioma syndrome	3232	Deafness-ear malformation-facial palsy syndrome
94087	Cytophagic histiocytic panniculitis	1566	Dandy-Walker malformation-postaxial polydactyly syndrome	3220	Deafness-enamel hypoplasia-nail defects syndrome
477787	Cytosolic phospholipase-A2 alpha deficiency associated bleeding disorder	2091	Daneman-Davy-Mancer syndrome	254898	Deafness-encephaloneuropathy-obesity-valvulopathy syndrome
		34587	Danon disease		
137678	Czech dysplasia, metatarsal type	99645	Dappled diaphyseal dysplasia	3218	Deafness-epiphyseal dysplasia-short stature syndrome
2736	Czeizel syndrome	218	Darier disease		
2917	Czeizel-Brooser syndrome	316	Darier-Gottron disease	3224	Deafness-genital anomalies-metacarpal and metatarsal synostosis syndrome
2437	Czeizel-Lozonci syndrome	218	Darier-White disease		
2953	D4ST1-deficient EDS	390	Darling disease	3237	Deafness-Hermann type symphalangism syndrome
2953	D4ST1-deficient Ehlers-Danlos syndrome	293978	DAVID syndrome		
90038	D+HUS	75565	Davies disease	90646	Deafness-hypogonadism syndrome
356978	D,L-2-HGA	2806	Dawson's encephalitis	94064	Deafness-infertility syndrome
356978	D,L-2-hydroxyglutaric acidemia	2143	DBS/FOAR syndrome	85321	Deafness-intellectual disability syndrome, Martin-Probst type
356978	D,L-2-hydroxyglutaric aciduria	1775	DC		
79315	D-2-HGA	79456	DCM	3226	Deafness-lymphedema-leukemia syndrome
79315	D-2-hydroxyglutaric acidemia	66634	DCMA syndrome		
79315	D-2-hydroxyglutaric aciduria	75381	DCMD	2408	Deafness-nephritis-ano-rectal malformation syndrome
93599	D-glycerate dehydrogenase deficiency	1653	DD		
941	D-glycerate kinase deficiency	99789	DD-I	3230	Deafness-oligodontia syndrome
941	D-glyceric acidemia	99791	DD-II	79500	Deafness-onychodystrophy-osteodystrophy-intellectual disability syndrome
941	D-glyceric aciduria	→231568	DDEB, Cockayne-Touraine type		
2134	D-HUS	231568	DDEB, generalized	79500	Deafness-onychodystrophy-osteodystrophy-intellectual disability-seizures syndrome
93581	D-HUS with anti-factor H antibodies	231568	DDEB, Pasini and Cockayne-Touraine types		
93578	D-HUS with B factor anomaly	231568	DDEB, Pasini type	79500	Deafness-onychoosteodystrophy-intellectual disability syndrome
93575	D-HUS with C3 anomaly	99970	DDEB-gen	79500	Deafness-onychoosteodystrophy-intellectual disability syndrome
357008	D-HUS with DGKE deficiency	99970	DDLS	→52368	Deafness-opticoacoustic nerve atrophy-dementia syndrome
93579	D-HUS with H factor anomaly	79499	DDOD syndrome		
93580	D-HUS with I factor anomaly	52368	DDON syndrome	123	Deafness-pili torti-hypogonadism syndrome
93576	D-HUS with MCP/CD46 anomaly	300536	DDOST-CDG	3219	Deafness-skeletal dysplasia-coarse face with full lips syndrome
217023	D-HUS with thrombomodulin anomaly	2962	De Bary syndrome		
1146	DA1	1130	De Die-Smulders-Vles-Fryns syndrome	3219	Deafness-skeletal dysplasia-lip granuloma syndrome
1146	DA1A	1598	De Grouchy syndrome		
		→782	De Hauwere syndrome	3217	Deafness-small bowel diverticulosis-neuropathy syndrome
		→782	De Hauwere-Chitty syndrome		

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3221	Deafness-thyroid hormone resistance syndrome	1617	Del(2)(q24)	261279	Del(17)(q23.1q23.2)
		251014	Del(2)(q31.1)	254346	Del(19)(p13.12)
3239	Deafness-vitiligo-achalasia syndrome	251019	Del(2)(q32)	357001	Del(19)(p13.13)
		251019	Del(2)(q32q33)	217346	Del(19)(q13.11)
→2697	Deal-Barrat-Dillon syndrome	251028	Del(2)(q33.1)	261295	Del(20)(p12.3)
158673	DEB, acral	1001	Del(2)(q37)	313781	Del(20)(p13)
79411	DEB, bullous dermolysis of the newborn	1621	Del(3)(q13)	444051	Del(20)(q11.2)
		356947	Del(3)(q26q27)	261311	Del(20)(q13.33)
89843	DEB, pruriginosa	397695	Del(3)(q27.3)	261323	Del(21)(q22.11q22.12)
158673	DEB-ac	65286	Del(3)(q29)	268261	Del(21)(q22.13q22.2)
79411	DEB-BDN	435638	Del(3)p(25.3)	96123	Del(22)
158676	DEB-na	238750	Del(4)(q21)	261476	Del(X)(p21)
89843	DEB-Pr	228384	Del(5)(q14.3)	1643	Del(X)(p23)
79410	DEB-Pt	314655	Del(5)(q31.3)	3034	Delayed membranous cranial ossification
431361	DECR deficiency with hyperlysinemia	251046	Del(6)(p22)		Delayed speech-facial asymmetry-strabismus-ear lobe creases syndrome
99970	Dedifferentiated liposarcoma	171829	Del(6)(q16)	3038	Delayed speech-facial asymmetry-strabismus-ear lobe creases syndrome
397587	Deep dermatophytosis	251056	Del(6)(q25)		
31150	Defective adenosine triphosphate-binding cassette transporter A1	251061	Del(7)(q31)	456298	Deletion 1p35.2
		251066	Del(8)(p11.2)	1606	Deletion 1p36
75496	Defective biosynthesis of proteodermatan sulfate	251071	Del(8)(p23.1)	1606	Deletion 1pter
		284160	Del(8)(q21.11)	1001	Deletion 2q37
60	Deficiency in Alpa-1-proteinase inhibitor	2496	Del(8)q(13)	1001	Deletion 2q37-qter
		324313	Del(9)(p13)	281	Deletion 5p
293978	Deficiency in anterior pituitary function-variable immunodeficiency syndrome	352665	Del(9)(q21)	1627	Deletion 5q35
		401923	Del(9)(q31.1q31.3)	904	Deletion 7q11.23
		284169	Del(10)(p11.21p12.31)	284160	Deletion 8q21.11
169150	Deficiency of complement of terminal pathway	276413	Del(10)(q22.3q23.3)	502	Deletion 8q24.1
		893	Del(11)(p13)	284169	Deletion 10p11.21p12.31
404546	Deficiency of IL-36R antagonist	444002	Del(11)(q22.2q22.3)	276413	Deletion 10q22.3q23.3
404546	Deficiency of IL-36Ra	2308	Del(11)(q23.3)	893	Deletion 11p13
158	Deficiency of plasma-membrane carnitine transporter	2308	Del(11)(qter)	94063	Deletion 12q14
		313884	Del(12)(p12.1)	289513	Deletion 12q15q21.1
679	Degos disease	280325	Del(12)(p13.33)	1587	Deletion 13q14
315	Degos genodermatosis "en cocardes"	94063	Del(12)(q14)	1590	Deletion 13q32
		289513	Del(12)(q15)(q21.1)	1600	Deletion 18q
1578	Dehydratase deficiency	412035	Del(13)(q12.3)	96123	Deletion 22
3202	Dehydrated hereditary stomatocytosis	1587	Del(13)(q14)	1647	Delleman syndrome
		96168	Del(13)(q34)	1647	Delleman-Oorthuys syndrome
64748	Dejerine-Sottas syndrome	261120	Del(14)(q11.2)	79101	Delta1-pyrroline-5-carboxylate dehydrogenase deficiency
2318	Dekaban-Arima syndrome	261144	Del(14)(q12)		
1627	Del (5)(q35)	264200	Del(14)(q22q23)	35664	Delta-1-pyrroline 5-carboxylate synthetase deficiency
1627	Del (5)(qter)	401935	Del(14)(q24.1q24.3)		
401986	Del(1)(p31p32)	261183	Del(15)(q11.2)	231237	Delta-beta-thalassemia
456298	Del(1)(p35.2)	199318	Del(15)(q13.3)	219	Delta-sarcoglycanopathy
1606	Del(1)(p36)	261190	Del(15)(q14)	168782	Dementia infantilis
250989	Del(1)(q21)	94065	Del(15)(q24)	97353	Dementia pugilistica
250999	Del(1)(q41q42)	261211	Del(16)(p11.2p12.2)	283	Demodicidosis
238769	Del(1)(q44)	261236	Del(16)(p13.11)	283	Demodicosis
293948	Del(1)p(21.3)	352629	Del(16)(q24.1)	314451	Demons-Meigs syndrome
363680	Del(2)(p13.2)	261250	Del(16)(q24.3)	79134	DEND syndrome
261349	Del(2)(p15p16.1)	97685	Del(17)(q11)	86903	Dendritic cell sarcoma not otherwise specified
163693	Del(2)(p21)	261265	Del(17)(q12)		
369881	Del(2)(p21) without cystinuria	363958	Del(17)(q21.31)		
228402	Del(2)(q23.1)				

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228423	Dendritic cell, monocyte, B and NK lymphoid deficiency	36426	Dermatostomatitis, Stevens Johnson type	99989	Developmental delay-epilepsy-neonatal diabetes syndrome, intermediate form
99828	Dengue fever	1660	Dermo-odonto dysplasia		
99828	Dengue virus infection	79149	Dermochondrocorneal dystrophy	2101	Developmental delay-hypotonia-extremities hypertrophy syndrome
2109	Dennis-Fairhurst-Moore syndrome	141051	Dermoid cyst of the face		
93571	Dense deposit disease	141046	Dermoid cyst of the neck		
1652	Dent disease	99688	Dermotrichic syndrome	459061	Developmental delay-short stature-dysmorphic features-sparse hair syndrome
93622	Dent disease type 1	1916	DES embryofetopathy		
93623	Dent disease type 2	1916	DES syndrome	79107	Developmental malformations-deafness-dystonia syndrome
1652	Dent syndrome	1425	Desbuquois dysplasia		
		1425	Desbuquois syndrome	209908	Developmental verbal dyspraxia
2095	Dental and eye anomalies-patent ductus arteriosus-normal intelligence syndrome	163703	DESC syndrome	71211	Devic disease
		228123	Desert fever	→3464	Devriendt-Legius-Fryns syndrome
1077	Dental ankylosis	228123	Desert rheumatism	1014	Devriendt-Vandenbergh-Fryns syndrome
101	Dentatorubral pallidolusian atrophy	98909	Desmin-related myofibrillar myopathy	403	Dexamethasone-sensitive hypertension
101	Dentatorubropallidolusian atrophy				
1653	Dentin dysplasia	84132	Desmin-related myopathy with Mallory body-like inclusions	1666	Dextrocardia
314721	Dentin dysplasia type 1 with microdontia and shape anomalies	98909	Desminopathy	→244	Dextrocardia-bronchiectasis-sinusitis syndrome
		873	Desmoid tumor		
99789	Dentin dysplasia type I	873	Desmoid type fibromatosis	99828	DF
99791	Dentin dysplasia type II			383	DFNX2
99792	Dentin dysplasia-sclerotic bones syndrome	251940	Desmoplastic infantile astrocytoma/ganglioglioma	31112	DFSP
49042	Dentinogenesis imperfecta	83469	Desmoplastic small round cell tumor	49042	DGI
166260	Dentinogenesis imperfecta type 2			49042	DGI without OI
166265	Dentinogenesis imperfecta type 3	251863	Desmoplastic/nodular medulloblastoma	166260	DGI-2
				373	DGSX
49042	Dentinogenesis imperfecta without osteogenesis imperfecta	35107	Desmosterolosis	319651	DHFR deficiency
				139518	dHMN1
166260	Dentinogenesis imperfecta, Shields type 2	98852	Desquamative interstitial pneumonia	139525	dHMN2
		158014	Destombes-Rosaï-Dorfman disease	139547	dHMN3 and dHMN4
166265	Dentinogenesis imperfecta, Shields type 3			139536	dHMN5
		163703	Devastating epileptic encephalopathy in school-aged children	98920	dHMN6
				139589	dHMN7
71267	Dentinogenesis imperfecta-short stature-hearing loss-intellectual disability syndrome	313892	Developmental and speech delay due to SOX5 deficiency	357043	dHMN with upper motor neuron signs
77295	Dentoleukoencephalopathy				
220	Denys-Drash syndrome	79157	Developmental delay due to 2-methylbutyryl-CoA dehydrogenase deficiency	139552	dHMNJ
3177	Der Kaloustian-Jarudi-Khoury syndrome			75376	DHRD
				49042	DI
3270	Der Kaloustian-McIntosh-Silver syndrome	289307	Developmental delay due to ALDH6A1 deficiency	166260	DI-2
369950	Der(8)t(8;12)			251940	DIA/DIG
96170	Der(22)t(11;22) syndrome	289307	Developmental delay due to methylmalonate semialdehyde dehydrogenase deficiency	3463	Diabetes insipidus-diabetes mellitus-optic atrophy-deafness syndrome
36397	Dercum disease				
1656	Dermatitis herpetiformis	289307	Developmental delay due to MMSDH deficiency	3464	Diabetes-hypogonadism-deafness-intellectual disability syndrome
1266	Dermato-cardio-skeletal syndrome, Borrone type	329195	Developmental delay with ASD and gait instability	1926	Diabetic embryopathy
31112	Dermatofibrosarcoma protuberans			85446	Dialysis-related amyloidosis
1659	Dermatoleukodystrophy	329195	Developmental delay with autism spectrum disorder and gait instability	85446	Dialysis-related arthropathy
221	Dermatomyositis			275523	Dianzani autoimmune lymphoproliferative disease
1657	Dermatoosteolysis, Kirghizian type	163988	Developmental delay-deafness syndrome, Hildebrand type	66637	Diaphanospondylodysostosis
86920	Dermatopathia pigmentosa reticularis	79134	Developmental delay-epilepsy-neonatal diabetes syndrome	255182	Diaphorase deficiency
				2141	Diaphragmatic defect-limb deficiency-skull defect syndrome

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2059	Diaphragmatic hernia-abnormal face-distal limb anomalies syndrome	220393	Diffuse cutaneous systemic scleroderma	973	Digits 2-5 oligodactyly, unilateral
				319651	Dihydrofolate reductase deficiency
2143	Diaphragmatic hernia-exomphalos-hypertelorism syndrome	220393	Diffuse cutaneous systemic sclerosis	79244	Dihydrolipoamide acetyltransferase component of pyruvate dehydrogenase complex deficiency
		2199	Diffuse erythrodermic palmoplantar keratoderma, Voerner type		
2143	Diaphragmatic hernia-hypertelorism-myopia-deafness syndrome	2199	Diffuse erythrodermic palmoplantar keratoderma, Vörner type	2394	Dihydrolipoamide dehydrogenase deficiency
98920	Diaphragmatic spinal muscular atrophy	702	Diffuse familial brain sclerosis	255182	Dihydrolipoyl dehydrogenase deficiency
		3165	Diffuse fasciitis with eosinophilia		
404521	Diaphragmatic spinal muscular atrophy type 2	300849	Diffuse large B-cell lymphoma of the central nervous system	79244	Dihydrolipoyllysine-residue acetyltransferase component of pyruvate dehydrogenase complex deficiency
1802	Diaphyseal dysplasia-anemia syndrome	300888	Diffuse large B-cell lymphoma with chronic inflammation		
85182	Diaphyseal medullary stenosis-bone malignancy syndrome	252031	Diffuse leptomeningeal melanocytosis	226	Dihydropteridine reductase deficiency
85182	Diaphyseal medullary stenosis-malignant fibrous histiocytoma syndrome	141209	Diffuse lymphangioma	38874	Dihydropyrimidinase deficiency
		141209	Diffuse lymphangiomatosis	1675	Dihydropyrimidine dehydrogenase deficiency
		141209	Diffuse lymphatic malformation		
103909	Diarrhea-vomiting due to trehalase deficiency	168811	Diffuse malignant peritoneal mesothelioma	38874	Dihydropyrimidinuria
97282	Diarrheogenic islet cell tumor	2123	Diffuse neonatal hemangiomatosis	99102	Dilatation of the left appendage
1671	Diastematomyelia			99102	Dilatation of the left auricle
628	Diastrophic dwarfism	86918	Diffuse palmoplantar hyperkeratosis-acrocyanosis syndrome	99101	Dilatation of the right atrial appendage
628	Diastrophic dysplasia			99101	Dilatation of the right atrial auricle
276603	Diazoxide-resistant focal hyperinsulinism due to Kir6.2 deficiency	369999	Diffuse palmoplantar keratoderma with painful fissures	66634	Dilated cardiomyopathy with ataxia
		2337	Diffuse palmoplantar keratoderma, Bothnian type	2229	Dilated cardiomyopathy-hypergonadotropic hypogonadism syndrome
276598	Diazoxide-resistant focal hyperinsulinism due to SUR1 deficiency	86918	Diffuse palmoplantar keratoderma-acrocyanosis syndrome	231111	DILE
2195	Dicarboxylic aminoaciduria	171700	Diffuse panbronchiolitis	243343	Dimethylglycine dehydrogenase deficiency
284343	DICER1 syndrome	71274	Diffuse peritoneal leiomyomatosis	→3157	Dincsoy-Salih-Patel syndrome
180086	Didelphys uterus	66627	Diffuse-type GCT	314002	Dinno syndrome
3463	DIDMOAD syndrome	66627	Diffuse-type giant cell tumor	1493	Dionisi-Vici-Sabetta-Gambarara syndrome
370046	Didymosis aplasticosebacea	567	DiGeorge sequence		
1672	Diencephalic cachexia	567	DiGeorge syndrome	227	Diphallia
1672	Diencephalic syndrome	238	Digestive duplication	1679	Diphtheria
1672	Diencephalic syndrome of childhood	141071	Digestive duplication cyst of the tongue	128	Diphyllbothriasis
1672	Diencephalic syndrome of emaciation			1681	Diprosopia
319192	Diencephalic-mesencephalic junction dysplasia	1305	Digital anomalies with short palpebral fissures and atresia of esophagus or duodenum	1681	Diprosopus
				1756	Dipygus
1916	Diethylstilbestrol embryofetopathy	391641	Digital anomalies with short palpebral fissures and atresia of esophagus or duodenum type 1	210115	DIRA
1916	Diethylstilbestrol syndrome			166291	Dirofilariasis
146	Differentiated thyroid carcinoma	352487	Digital anomalies-intellectual disability-short stature syndrome	94064	DIS
90060	Diffuse alveolar hemorrhage			35122	Disaccharide intolerance
324	Diffuse angiokeratoma	31828	Digitalis poisoning	306446	Disaccharide intolerance with minimal starch tolerance
251595	Diffuse astrocytoma				
404437	Diffuse cerebral and cerebellar atrophy-intractable seizures-progressive microcephaly syndrome	1146	Digitotalar dysmorphism	306474	Disaccharide intolerance with starch and lactose intolerance
		294990	Digits 2-5 hypodactyly	306436	Disaccharide intolerance with starch intolerance
		295114	Digits 2-5 hypodactyly, bilateral	306462	Disaccharide intolerance without starch intolerance
79456	Diffuse cutaneous maculopapulous mastocytosis	973	Digits 2-5 hypodactyly, unilateral		
79456	Diffuse cutaneous mastocytosis	294990	Digits 2-5 oligodactyly	306486	Disaccharide intolerance without sucrose intolerance
		295114	Digits 2-5 oligodactyly, bilateral		

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90281	Discoid lupus erythematosus	178400	Distal anterior compartment myopathy	96098	Distal duplication 6q
216694	Discordant ventriculoarterial and atrioventricular connections		1146	Distal arthrogryposis type 1	96074
99052	Discrete fibromuscular subaortic stenosis	2053	Distal arthrogryposis type 2A	96100	Distal duplication 8q
		1147	Distal arthrogryposis type 2B	96101	Distal duplication 9q
99051	Discrete fixed membranous subaortic stenosis	376	Distal arthrogryposis type 3	96102	Distal duplication 10q
		65720	Distal arthrogryposis type 4	96103	Distal duplication 11q
90394	Discrete papular lichen myxedematosus	1154	Distal arthrogryposis type 5	96105	Distal duplication 13q
139420	Disease-associated transverse myelitis	329457	Distal arthrogryposis type 5 without ophthalmoparesis	1705	Distal duplication 14q
		329457	Distal arthrogryposis type 5 without ophthalmoplegia	1707	Distal duplication 15q
210272	Disembarkment syndrome	329457	Distal arthrogryposis type 5D	96078	Distal duplication 16p
2412	Dislocation of the hip-dysmorphism syndrome		1144	Distal arthrogryposis type 6	96106
8	Disomy Y	3377	Distal arthrogryposis type 7	3379	Distal duplication 17q
2983	Disorder of sex development-intellectual disability syndrome	65743	Distal arthrogryposis type 8	1716	Distal duplication 18q
		115	Distal arthrogryposis type 9	1717	Distal duplication 19q
345	Dissecting cellulitis of the scalp	251515	Distal arthrogryposis type 10	96107	Distal duplication 20q
54251	Disseminated aseptic abscesses	376	Distal arthrogryposis type IIA	96109	Distal duplication 22q
1306	Disseminated dermatofibrosis with osteopoikilosis	1154	Distal arthrogryposis type IIB	1762	Distal duplication Xq
		65720	Distal arthrogryposis type IID	139518	Distal hereditary motor neuropathy type 1
397587	Disseminated granulomatous dermatophytosis	1154	Distal arthrogryposis with ophthalmoplegia	139525	Distal hereditary motor neuropathy type 2
141209	Disseminated lymphangioma	254351	Distal del(7)(q11.23)	139547	Distal hereditary motor neuropathy type 3 and type 4
141209	Disseminated lymphangiomatosis		261222	Distal del(16)(p11.2)	139536
141209	Disseminated lymphatic malformation	319171	Distal del(17)(p13.1)	98920	Distal hereditary motor neuropathy type 6
228264	Disseminated nevus anelasticus	261257	Distal del(17)(p13.3)		139589
71274	Disseminated peritoneal leiomyomatosis	261330	Distal del(22)(q11.2)	139536	Distal hereditary motor neuropathy type V
		36367	Distal deletion 1q		357043
79152	Disseminated superficial actinic porokeratosis	280	Distal deletion 4p	139552	Distal hereditary motor neuropathy, Jerash type
1620	Distal 3p deletion	96145	Distal deletion 4q		139536
1627	Distal 5q deletion	96125	Distal deletion 6p	1307	Distal limb deficiencies-micrognathia syndrome
254351	Distal 7q11.23 microdeletion syndrome	96126	Distal deletion 7p		36367
		1636	Distal deletion 7q36	1620	Distal monosomy 3p
261102	Distal 7q11.23 microduplication syndrome	1642	Distal deletion 9p	280	Distal monosomy 4p
		96148	Distal deletion 10q	96145	Distal monosomy 4q
1580	Distal 10p deletion	2308	Distal deletion 11q	96125	Distal monosomy 6p
1590	Distal 13q deletion	280325	Distal deletion 12p	96126	Distal monosomy 7p
1596	Distal 15q deletion syndrome	96149	Distal deletion 12q	254351	Distal monosomy 7q11.23
		96168	Distal deletion 13q34	1636	Distal monosomy 7q36
261222	Distal 16p11.2 microdeletion syndrome	96150	Distal deletion 14q	1642	Distal monosomy 9p
		96129	Distal deletion 19p	1580	Distal monosomy 10p
319171	Distal 17p13.1 microdeletion syndrome	96152	Distal deletion 20q	96148	Distal monosomy 10q
		261102	Distal dup(7)(q11.23)	2308	Distal monosomy 11q
261257	Distal 17p13.3 microdeletion syndrome	261337	Distal dup(22)(q11.2)	280325	Distal monosomy 12p
		293939	Distal dup(X)q(28)	96149	Distal monosomy 12q
1597	Distal 17q deletion	96069	Distal duplication 1p36	1590	Distal monosomy 13q
261330	Distal 22q11.2 microdeletion syndrome	96070	Distal duplication 2p	96150	Distal monosomy 14q
		96094	Distal duplication 2q		
261337	Distal 22q11.2 microduplication syndrome	96071	Distal duplication 3p		
		96072	Distal duplication 4p		
63273	Distal ABD-filaminopathy	96096	Distal duplication 4q		
399096	Distal anoctaminopathy	96097	Distal duplication 5q		
		1745	Distal duplication 6p		

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1596	Distal monosomy 15q	96101	Distal trisomy 9q	86309	Dolichyl-phosphate N-acetylgalactosamine phosphotransferase deficiency
261222	Distal monosomy 16p11.2	96102	Distal trisomy 10q		
261257	Distal monosomy 17p13.3	96103	Distal trisomy 11q	3427	DOLV
1597	Distal monosomy 17q	96105	Distal trisomy 13q	231226	Dominant beta-thalassemia
96129	Distal monosomy 19p13.3	1705	Distal trisomy 14q	75376	Dominant drusen
96152	Distal monosomy 20q	1707	Distal trisomy 15q	898	Dominant hyaloideoretinal dystrophy of Wagner
261330	Distal monosomy 22q11.2	96078	Distal trisomy 16p	244305	Dominant hypophosphatemia with nephrolithiasis or osteoporosis
59135	Distal myopathy type 1	96106	Distal trisomy 16q	276580	Dominant KATP hyperinsulinism due to Kir6.2 deficiency
399086	Distal myopathy type 3	3379	Distal trisomy 17q	75376	Dominant radial drusen
178400	Distal myopathy with anterior tibial onset	1716	Distal trisomy 18q	90035	Donath-Landsteiner hemolytic anemia
		1717	Distal trisomy 19q	90035	Donath-Landsteiner syndrome
34521	Distal myopathy with early respiratory muscle involvement	96107	Distal trisomy 20q	2143	Donnai-Barrow syndrome
		96109	Distal trisomy 22q	508	Donohue syndrome
63273	Distal myopathy with posterior leg and anterior hand involvement	261337	Distal trisomy 22q11.2	79500	DOOR syndrome
		293939	Distal trisomy Xq28	79500	DOORS syndrome
602	Distal myopathy with rimmed vacuoles	293939	Distal Xq28 microduplication syndrome	1942	Doose syndrome
600	Distal myopathy with vocal cord weakness	98920	Distal-HMN type 6	70594	Dopa-responsive dystonia due to sepiapterin reductase deficiency
602	Distal myopathy, Nonaka type	→33001	Distichiasis-congenital heart defects-peripheral vascular anomalies syndrome		
609	Distal myopathy, Udd type			1916	Distilbene embryofetopathy
603	Distal myopathy, Welander type	1685	Distomatosis		
98911	Distal myotilinopathy	1685	Distomiasis	98907	Dorfman-Chanarin disease
2776	Distal osteolysis-short stature-intellectual disability syndrome	404546	DITRA	3426	DORV
18	Distal renal tubular acidosis	99099	Divided left atrium	423712	DORV with atrioventricular septal defect, pulmonary stenosis, heterotaxy
→402041	Distal renal tubular acidosis type 1b	99098	Divided right atrium		
→402041	Distal renal tubular acidosis type 1c	91131	DK1-CDG	99046	DORV with non-committed subpulmonary VSD
93610	Distal renal tubular acidosis with anemia	3439	DK phocomelia syndrome		
		1775	DKC	423693	DORV with subaortic or doubly committed VSD
139525	Distal spinal muscular atrophy type 2	300849	DLBCL of the CNS		
139547	Distal spinal muscular atrophy type 3	300888	DLBCL with chronic inflammation	99043	DORV with subaortic or doubly committed VSD with pulmonary stenosis
		2394	DLD deficiency		
206580	Distal spinal muscular atrophy type 4	252031	DLM	99045	DORV with subpulmonary VSD
		221	DM		
139536	Distal spinal muscular atrophy type 5	273	DM1	99043	DORV, Fallot type
		98896	DMD	99045	DORV-TGA
139589	Distal spinal muscular atrophy with vocal cord paralysis	243343	DMG dehydrogenase deficiency	869	Double A syndrome
3248	Distal symphalangism	602	DMRV	216694	Double discordance
314588	Distal tetrasomy 15q	99812	DNA ligase IV deficiency	1464	Double inlet left ventricle
609	Distal titinopathy	251946	DNET	141091	Double nose
96069	Distal trisomy 1p36	404443	DNMT3A-related overgrowth syndrome	3427	Double outlet left ventricle
96070	Distal trisomy 2p	251975	DNT of the cerebellum	3426	Double outlet right ventricle
96094	Distal trisomy 2q	1215	DOA+	423712	Double outlet right ventricle with atrioventricular septal defect, pulmonary stenosis, heterotaxy
96071	Distal trisomy 3p	447737	DOCK2 deficiency		
96072	Distal trisomy 4p	217390	DOCK8 immunodeficiency syndrome	→423693	Double outlet right ventricle with doubly committed ventricular septal defect
96096	Distal trisomy 4q	79322	Dol-P-mannosyltransferase deficiency		
96097	Distal trisomy 5q	91131	Dolichol kinase deficiency	99046	Double outlet right ventricle with non-committed subpulmonary ventricular septal defect
1745	Distal trisomy 6p	2616	Dolichospondylic dysplasia		
96098	Distal trisomy 6q				
96074	Distal trisomy 7p				
261102	Distal trisomy 7q11.23				
96100	Distal trisomy 8q				

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423693	Double outlet right ventricle with subaortic or doubly committed ventricular septal defect	90037	Drug-induced autoimmune hemolytic anemia	228399	Dup(8)(q12)
				276422	Dup(10)(q22.3q23.3)
99043	Double outlet right ventricle with subaortic or doubly committed ventricular septal defect with pulmonary stenosis	90157	Drug-induced localized lipodystrophy	300305	Dup(11)p(15.4)
		231111	Drug-induced lupus erythematosus	261229	Dup(14)(q11.2)
		464453	Drug-induced methemoglobinemia	238446	Dup(15)(q11q13)
		251325	Drug-induced vasculitis	261204	Dup(16)(p11.2p12.2)
→423693	Double outlet right ventricle with subaortic ventricular septal defect	97368	Drug-related renal tubular dysgenesis	261243	Dup(16)(p13.11)
99045	Double outlet right ventricle with subpulmonary ventricular septal defect	94086	Drummond syndrome	96078	Dup(16)(p13.3)
		33069	DS	477817	Dup(17)(p11.2p12)
		99887	DS-AMKL	217385	Dup(17)(p13.3)
99045	Double outlet right ventricle with transposition of the great arteries	98920	dSMA1	139474	Dup(17)(q11.2)
		139525	dSMA2	261272	Dup(17)(q12)
99043	Double outlet right ventricle, Fallot type	139547	dSMA3	217340	Dup(17)(q21.31)
		206580	dSMA4	261290	Dup(17p)
3286	Double tachycardia induced by catecholamines	314485	dSMA5	447980	Dup(19)(p13.13)
3411	Double uterus and obstructed hemivagina syndrome	139557	DSMAX	363659	Dup(20)(q11.2)
		83469	DSRCT	261318	Dup(20p)
3411	Double uterus-hemivagina-renal agenesis syndrome	412181	DST-related epidermolysis bullosa simplex	1727	Dup(22)(q11)
8	Double Y	99789	DTDP1	284180	Dup(X)(p22)
95474	Double-orifice mitral valve	99791	DTDP2	284180	Dup(X)(p22.13p22.2)
79145	Dowling-Degos disease	2639	Du Pan syndrome	314389	Dup(X)(q12-q13.3)
870	Down syndrome	50817	Duane anomaly-myopathy-scoliosis syndrome	261483	Dup(X)(q27.3q28)
75376	Doyme honeycomb retinal dystrophy	233	Duane retraction syndrome	261344	Duplication 1q
		233	Duane syndrome	1738	Duplication 4p
86309	DPAGT1-CDG	93293	Duane-radial ray syndrome	1742	Duplication 5p
314621	DPG-plus syndrome	261647	Duane-radial ray syndrome due to a point mutation	264450	Duplication 8p
71274	DPL	261638	Duane-radial ray syndrome due to monosomy 20q13	1752	Duplication 8q
79322	DPM1-CDG			96167	Duplication 8q/deletion 8p
329178	DPM2-CDG	234	Dubin-Johnson syndrome	236	Duplication 9p
263494	DPM3-CDG	234	Dubin-Sprinz disease	1699	Duplication 12p
231	Dracunculiasis	235	Dubowitz syndrome	1715	Duplication 18p
231	Dracunculosis	98896	Duchenne muscular dystrophy	1727	Duplication 22q11.2
220	Drash syndrome	280315	Duct-centric pancreatitis	261318	Duplication of 20p
33069	Dravet syndrome	2442	Duncan disease	314621	Duplication of the pituitary gland
→79500	DRC syndrome	2348	Dunnigan syndrome	314621	Duplication of the pituitary gland-plus syndrome
70594	DRD due to SRD	→293864	Duodenal and extrahepatic biliary atresia-hypoplastic pancreas-intestinal malrotation syndrome	1738	Duplication of the short arm of chromosome 4
130	Dream disease	1203	Duodenal atresia	1742	Duplication of the short arm of chromosome 5
139402	DRESS syndrome	250994	Dup(1)(q21.1)	236	Duplication of the short arm of chromosome 9
101	DRPLA	313947	Dup(2)(q23.1)	1715	Duplication of the short arm of chromosome 18
233	DRS	294026	Dup(2)(q31.1)	237	Duplication of urethra
18	dRTA	96095	Dup(3)(q26)	284180	Duplication Xp22
→402041	dRTA type 1b	329802	Dup(5)(p13)	3306	Duplication/inversion 15q11
→402041	dRTA type 1c	228415	Dup(5)(q35)	97339	Dural sinus malformation
93610	dRTA with anemia	314034	Dup(7)(p22.1)	1656	Durhing-Brocq disease
264978	Drug or radiation exposure-related interstitial lung disease	96121	Dup(7)(q11.23)	233	DURS
		459074	Dup(7)(q36.3)	→331176	Dursun syndrome
		251076	Dup(8)(p23.1p23.1)	98984	Dusty cataract
139402	Drug rash with eosinophilia and systemic symptoms			3377	Dutch-Kentucky syndrome
139402	Drug reaction eosinophilic systemic syndrome				
90037	Drug-induced AIHA				

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→969	Dwarfism-stiff joint-ocular abnormalities syndrome	2204	Dysplastic cortical hyperostosis	1934	Early infantile epileptic encephalopathy
→2616	Dwarfism-tall vertebrae syndrome	65285	Dysplastic gangliocytoma of the cerebellum		Early infantile epileptic encephalopathy with suppression-bursts
1566	DWM with postaxial polydactyly	325	Dysprothrombinemia	1934	Early infantile epileptic encephalopathy without suppression burst
239	Dyggve-Melchior-Clausen disease	2476	Dysraphism-cleft lip/palate-limb reduction defects syndrome	369894	Early infantile epileptic encephalopathy with suppression-bursts
2274	Dykes-Markes-Harper syndrome		Dyssegmental dysplasia, Rolland-Desbuquois type	1935	Early infantile epileptic encephalopathy without suppression burst
464306	DYRK1A-related intellectual disability syndrome	156731	Dyssegmental dysplasia, Silverman-Handmaker type	1935	Early myoclonic encephalopathy
268261	DYRK1A-related intellectual disability syndrome due to 21q22.13q22.2 microdeletion	1865	Dyssegmental dysplasia-glaucoma syndrome	1935	Early myoclonic encephalopathy with suppression-bursts
		1804	Dyssegmental dysplasia-glaucoma syndrome	98988	Early-onset anterior polar cataract
464311	DYRK1A-related intellectual disability syndrome due to a point mutation	85198	Dysspondyloenchondromatosis	98988	Early-onset anterior subcapsular cataract
		71517	Dystonia 12		Early-onset anterior subcapsular cataract
296	Dyschondroplasia	→98808	Dystonia 14	1020	Early-onset autosomal dominant Alzheimer disease
1765	Dyschondrosteosis-nephritis syndrome	210571	Dystonia 16		Early-onset benign childhood occipital epilepsy
		98811	Dystonia 18	98815	Early-onset benign childhood occipital epilepsy
41	Dyschromatosis symmetrica hereditaria	420492	Dystonia 23		Early-onset cataract with Y-shaped suture opacities
		420485	Dystonia 24	98985	Early-onset cataract with Y-shaped suture opacities
241	Dyschromatosis universalis	256	Dystonia musculorum deformans		Early-onset cerebellar ataxia with retained tendon reflexes
251946	Dysembryoplastic neuroepithelial tumor	412217	Dystonia-aphonia syndrome	1177	Early-onset cerebellar ataxia with retained tendon reflexes
251975	Dysembryoplastic neuroepithelial tumor of cerebellum	199351	Dystonia-parkinsonism, Paisan-Ruiz type	84132	Early-onset desmin-related myopathy
1766	Dysequilibrium syndrome	293381	Dystrophia Helsinglandica		Early-onset desmin-related myopathy
99912	Dyserginomatous germ cell cancer of ovary	293381	Dystrophia Smolandiensis	1667	Early-onset diabetes mellitus with multiple epiphyseal dysplasia
		79409	Dystrophic epidermolysis bullosa inversa	210571	Early-onset diabetes mellitus with multiple epiphyseal dysplasia
3010	Dysharmonic skeletal maturation-muscular fiber disproportion syndrome	89843	Dystrophic epidermolysis bullosa pruriginosa	289266	Early-onset dystonia parkinsonism
1775	Dyskeratosis congenita	158676	Dystrophic epidermolysis bullosa, nails only		Early-onset epileptic encephalopathy and intellectual disability due to GRIN2A mutation
3088	Dyskeratosis congenita with bilateral exudative retinopathy	256	DYT1	411986	Early-onset epileptic encephalopathy-cortical blindness-intellectual disability-facial dysmorphism syndrome
412	Dyslipidemia type 3	99657	DYT2		Early-onset epileptic encephalopathy-cortical blindness-intellectual disability-facial dysmorphism syndrome
1779	Dysmorphism-cleft palate-loose skin syndrome	53351	DYT3	1020	Early-onset familial autosomal dominant Alzheimer disease
		98805	DYT4		Early-onset familial autosomal dominant Alzheimer disease
289553	Dysmorphism-conductive hearing loss-heart defect syndrome	98808	DYT5a	256	Early-onset generalized limb-onset dystonia
		101150	DYT5b		Early-onset generalized limb-onset dystonia
1780	Dysmorphism-multiple structural anomalies syndrome	98806	DYT6	256	Early-onset generalized limb-onset dystonia
		53583	DYT9		Early-onset generalized limb-onset dystonia
2104	Dysmorphism-pectus carinatum-joint laxity syndrome	71517	DYT12	88660	Early-onset generalized limb-onset dystonia
		98807	DYT13		Early-onset generalized limb-onset dystonia
2282	Dysmorphism-short stature-deafness-disorder of sex development syndrome	→98808	DYT14	324290	Early-onset hypertension with exacerbation in pregnancy
		→36899	DYT15	441452	Early-onset Lafora body disease
		210571	DYT16		Early-onset lamellar cataract
2282	Dysmorphism-short stature-deafness-pseudohermaphroditism syndrome	98811	DYT18	79242	Early-onset multiple carboxylase deficiency
		306734	DYT21		Early-onset multiple carboxylase deficiency
		420492	DYT23	289377	Early-onset myopathy with fatal cardiomyopathy
1782	Dysosteosclerosis	420485	DYT24		Early-onset myopathy with fatal cardiomyopathy
800	Dysostosis enchondralis metaepiphysaria, Catel-Hempel type	2394	E3-deficient maple syrup urine disease	439212	Early-onset myopathy-areflexia-respiratory distress-dysphagia syndrome
1798	Dysostosis, Stanescu type	231249	E-beta-thalassemia	91492	Early-onset myopathy-areflexia-respiratory distress-dysphagia syndrome
99082	Dysphagia lusoria	2970	Eagle-Barret syndrome	98991	Early-onset non-syndromic cataract
1822	Dysplasia epiphysealis hemimelica	40923	Eales disease	2828	Early-onset nuclear cataract
168621	Dysplasia of head of femur, Meyer type	2554	Ear-patella-short stature syndrome	2379	Early-onset Parkinson disease
					Early-onset parkinsonism-intellectual disability syndrome

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
98992	Early-onset partial cataract	→1896	ECP syndrome	1896	Ectrodactyly-ectodermal dysplasia-cleft lip/palate syndrome
98993	Early-onset posterior polar cataract	99102	Ectasia of the left appendage	1892	Ectrodactyly-polydactyly syndrome
441447	Early-onset posterior subcapsular cataract	99102	Ectasia of the left auricle	1894	Ectrodactyly-spina bifida-cardiopathy syndrome
256	Early-onset primary dystonia	99101	Ectasia of the right atrial appendage	1997	Ectropion inferior-cleft lip and or palate syndrome
157941	Early-onset prion disease with prominent psychiatric features	99101	Ectasia of the right atrial auricle	906	Eczema-thrombocytopenia-immunodeficiency syndrome
352654	Early-onset progressive neurodegeneration-blindness-ataxia-spasticity syndrome	35737	Ectasic coloboma	98813	EDA-ID
→90340	Early-onset progressive neurodegeneration-blindness-ataxia-spasticity syndrome	→1071	Ectodermal dysplasia syndrome, Rapp-Hodgkin type	247827	EDCS
96369	Early-onset schizophrenia	69083	Ectodermal dysplasia with natal teeth, Turnpenny type	293936	EDICT syndrome
364055	Early-onset severe retinal dystrophy	1809	Ectodermal dysplasia with skin anomalies and intellectual disability	1895	Edinburgh malformation syndrome
313772	Early-onset spastic ataxia-neuropathy syndrome	1816	Ectodermal dysplasia, Berlin type	93308	EDM1
98985	Early-onset sutural cataract	→1071	Ectodermal dysplasia, Rapp-Hodgkin type	93307	EDM4
256	Early-onset torsion dystonia	1818	Ectodermal dysplasia, trichoodontoonychial type	93311	EDM5
1243	Early-onset vitelliform macular dystrophy	→1658	Ectodermal dysplasia-absent dermatoglyphs syndrome	261	EDMD
98890	Early-onset X-linked optic atrophy	140936	Ectodermal dysplasia-absent dermatoglyphs syndrome	98863	EDMD1
98995	Early-onset zonular cataract	→2036	Ectodermal dysplasia-acanthosis nigricans syndrome	98853	EDMD2
199343	EAST syndrome	1806	Ectodermal dysplasia-adrenal cyst syndrome	98855	EDMD3
391320	East Texas bleeding disorder	3354	Ectodermal dysplasia-adrenal cyst syndrome	90309	EDS I
83594	Eastern equine encephalitis	247827	Ectodermal dysplasia-adrenal cyst syndrome	90318	EDS II
83594	Eastern equine encephalomyelitis	1806	Ectodermal dysplasia-blindness syndrome	285	EDS III
1973	Eastman-Bixler syndrome	3354	Ectodermal dysplasia-blindness syndrome	286	EDS IV
166418	Eating seizures	247827	Ectodermal dysplasia-cataracts-kyphoscoliosis syndrome	198	EDS IX
86880	EATL	1897	Ectodermal dysplasia-cataracts-kyphoscoliosis syndrome	286	EDS type 4
79406	EB progressive	1812	Ectodermal dysplasia-cutaneous syndactyly syndrome	75497	EDS V
79405	EBJ-I	1883	Ectodermal dysplasia-cutaneous syndactyly syndrome	1900	EDS VIA
319218	Ebola fever	423454	Ectodermal dysplasia-ectrodactyly-macular dystrophy syndrome	1899	EDS VII
319218	Ebola hemorrhagic fever	158668	Ectodermal dysplasia-ectrodactyly-macular dystrophy syndrome	99875	EDS VIIA
319218	Ebola virus disease	247820	Ectodermal dysplasia-intellectual disability-central nervous system malformation syndrome	99876	EDS VIIIB
412181	EBS-AR BP230	448270	Ectodermal dysplasia-intellectual disability-central nervous system malformation syndrome	1901	EDS VIIC
412189	EBS-AR exophilin 5	1885	Ectodermal dysplasia-intellectual disability-central nervous system malformation syndrome	75392	EDS VIII
89838	EBS-AR KRT14	1884	Ectodermal dysplasia-sensorineural deafness syndrome	82004	EDS with periventricular heterotopia
79400	EBS-loc	99889	Ectodermal dysplasia-sensorineural deafness syndrome	300179	EDS with progressive kyphoscoliosis, myopathy, and deafness
257	EBS-MD	231632	Ectodermal dysplasia-short stature syndrome	300179	EDS with progressive kyphoscoliosis, myopathy, and hearing loss
158681	EBS-migr	99889	Ectodermal dysplasia-short stature syndrome	75501	EDS X
79397	EBS-MP	95496	Ectodermal dysplasia-skin fragility syndrome	2295	EDS XI
79401	EBS-O	2440	Ectodermal dysplasia-skin fragility syndrome	2953	EDS, arthrogyposic type
158684	EBS-PA	→1896	Ectopia cordis	230851	EDS, cardiac valvular type
89839	EBSS	→1896	Ectopia lentis syndrome	287	EDS, classic type
1880	Ebstein anomaly of the tricuspid valve	→1896	Ectopia lentis-chorioretinal dystrophy-myopia syndrome	230839	EDS, classic-like type
1880	Ebstein malformation	→1896	Ectopia lentis-chorioretinal dystrophy-myopia syndrome	2953	EDS, Kosho type
313920	EBV-associated gastric carcinoma	99889	Ectopic ACTH secreting tumor	300179	EDS, kyphoscoliotic and hearing loss type
289661	EBV-positive DLBCL of the elderly	95496	Ectopic aldosterone-producing tumor	1900	EDS, kyphoscoliotic type
313920	EBVaGC	2440	Ectopic Cushing syndrome	2953	EDS, musculocontractural type
50944	Eccrine tumors-ectodermal dysplasia	→1896	Ectopic neurohypophysis	1900	EDS, oculoscoliotic type
284	Echinococcus multilocularis infection	→1896	Ectrodactyly	75496	EDS, progeroid type
199332	ECO syndrome	→1896	Ectrodactyly-cleft palate syndrome		
		→1896	Ectrodactyly-ectodermal dysplasia without clefting syndrome		

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157965	EDS, spondylocheirodysplastic type	230851	Ehlers-Danlos syndrome, cardiac valvular type	221054	Elejalde syndrome
230845	EDS, vascular-like type			289	Ellis Van Creveld syndrome
230857	EDS/OI syndrome	287	Ehlers-Danlos syndrome, classic type	2516	Ellis-Yale-Winter syndrome
247820	EDSS			1299	Elsahy-Waters syndrome
247820	EDSS1	230839	Ehlers-Danlos syndrome, classic-like type	1997	Elsching syndrome
247827	EDSS2			96170	Emanuel syndrome
178464	Edström Myopathy	1901	Ehlers-Danlos syndrome, dermatosparaxis type	439212	EMARDD
3380	Edwards syndrome			1942	EMAS
2668	Edwards-Patton-Dilly syndrome	75501	Ehlers-Danlos syndrome, fibronectin-deficient	3226	Emberger syndrome
322	EEC	75501	Ehlers-Danlos syndrome, fibronectinemic type	1914	Embryofetopathy due to oral anticoagulant therapy
1896	EEC syndrome			180226	Embryonal carcinoma
→1896	EEC syndrome without cleft lip/palate	285	Ehlers-Danlos syndrome, hypermobile type	48736	Embryonal carcinoma of the central nervous system
1897	EEM syndrome			48736	Embryonal carcinoma of the CNS
357131	Effort subclavian vein thrombosis	285	Ehlers-Danlos syndrome, hypermobility type	99757	Embryonal rhabdomyosarcoma
101039	EFMR			178315	Embryonal sarcoma of the liver
2070	EGE	2953	Ehlers-Danlos syndrome, Kosho type	1664	Embryonary disorganization syndrome
183	EGPA			983	Embryonic testicular regression syndrome
319218	EHF	300179	Ehlers-Danlos syndrome, kyphoscoliotic and deafness type	139431	EMEA
312	EHK	300179	Ehlers-Danlos syndrome, kyphoscoliotic and hearing loss type	98863	Emerinopathy
230839	Ehlers-Danlos syndrome due to tenascin-X deficiency	1900	Ehlers-Danlos syndrome, kyphoscoliotic type	261	Emery-Dreifuss muscular dystrophy
90309	Ehlers-Danlos syndrome type 1	2953	Ehlers-Danlos syndrome, musculocontractural type	1927	Emery-Nelson syndrome
90318	Ehlers-Danlos syndrome type 2			485418	EMILIN-1-related connective tissue disease
285	Ehlers-Danlos syndrome type 3	1900	Ehlers-Danlos syndrome, oculoscoliotic type	83600	Encephalitis lethargica
286	Ehlers-Danlos syndrome type 4			221126	Encephaloclastic proliferative vasculopathy
75497	Ehlers-Danlos syndrome type 5	75392	Ehlers-Danlos syndrome, periodontitis type	2396	Encephalocraniocutaneous lipomatosis
1900	Ehlers-Danlos syndrome type 6A			3205	Encephalofacial angiomatosis
1899	Ehlers-Danlos syndrome type 7	75496	Ehlers-Danlos syndrome, progeroid type	1035	Encephalopathy due to beta-mercaptolactate-cysteine disulfiduria
99875	Ehlers-Danlos syndrome type 7A			71277	Encephalopathy due to GLUT1 deficiency
99876	Ehlers-Danlos syndrome type 7B	157965	Ehlers-Danlos syndrome, spondylocheirodysplastic type	79155	Encephalopathy due to hydroxykynureninuria
1901	Ehlers-Danlos syndrome type 7C			139406	Encephalopathy due to prosaposin deficiency
75392	Ehlers-Danlos syndrome type 8	286	Ehlers-Danlos syndrome, vascular type	833	Encephalopathy due to sulfite oxidase deficiency
198	Ehlers-Danlos syndrome type 9			210128	Encephalopathy due to urocanase deficiency
75501	Ehlers-Danlos syndrome type 10	230845	Ehlers-Danlos syndrome, vascular-like type	51	Encephalopathy with basal ganglia calcification
2295	Ehlers-Danlos syndrome type 11			51	Encephalopathy with intracranial calcification and chronic lymphocytosis of cerebrospinal fluid
286	Ehlers-Danlos syndrome type IV	230857	Ehlers-Danlos/osteogenesis imperfecta syndrome	319678	Encephalopathy-hypertrophic cardiomyopathy-renal tubular disease syndrome
198	Ehlers-Danlos syndrome type IX				
82004	Ehlers-Danlos syndrome with periventricular heterotopia	1902	Ehrlichiosis		
75501	Ehlers-Danlos syndrome with platelet dysfunction from fibronectin abnormality	820	Ehrmann-Sneddon syndrome		
		312	EI		
300179	Ehlers-Danlos syndrome with progressive kyphoscoliosis, myopathy, and deafness	1934	EIEE		
		165991	EIHI		
		79106	Eiken syndrome		
300179	Ehlers-Danlos syndrome with progressive kyphoscoliosis, myopathy, and hearing loss	97214	Eisenmenger syndrome		
		476096	EKC syndrome		
		317	EKV		
1899	Ehlers-Danlos syndrome, arthrochalasia type	228240	Elastoderma		
		228243	Elastofibroma dorsi		
		228254	Elastoma		
1899	Ehlers-Danlos syndrome, arthrochalis type	79148	Elastosis perforans serpiginosa		
		228236	Elastotic striae		
2953	Ehlers-Danlos syndrome, arthrogryposic type	33445	Elejalde disease		

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1261	Encephalopathy-intracerebral calcification-retinal degeneration syndrome	402035	Eosinophilic colitis	79401	Epidermolysis bullosa simplex, Ogna type
		75566	Eosinophilic endocarditis		
		2070	Eosinophilic enteritis		
3205	Encephalotrigeminal angiomatosis	73247	Eosinophilic esophagitis	79400	Epidermolysis bullosa simplex, Weber-Cockayne type
296	Enchondromatosis	3165	Eosinophilic fasciitis	312	Epidermolytic hyperkeratosis
99075	Encircling double aortic arch	2070	Eosinophilic gastroenteritis	312	Epidermolytic ichthyosis
199332	Endocrine-cerebro-osteodysplasia syndrome	2070	Eosinophilic gastroenterocolitis	2199	Epidermolytic palmoplantar keratoderma
		99871	Eosinophilic granuloma		
876	Endodermal sinus tumor	183	Eosinophilic granulomatosis with polyangiitis	2199	Epidermolytic palmoplantar keratoderma of Voerner
252006	Endodermal sinus tumor of central nervous system				
252006	Endodermal sinus tumor of CNS	482	Eosinophilic lymphogranuloma	2199	Epidermolytic palmoplantar keratoderma of Vörner
98974	Endoepithelial corneal dystrophy	364055	EOSRD	141077	Epignathus
213741	Endometrial adenoid cystic carcinoma	256	EOTD		
		251880	Ependyoblastoma	65683	Epilepsy due to FCD
213726	Endometrial capillary carcinoma	251636	Ependymoma	86911	Epilepsy with myoclonic absences
		99169	Epiblepharon	1942	Epilepsy with myoclonic-astatic seizures
213716	Endometrial squamous cell carcinoma	185	Epibronchial right pulmonary vein syndrome		
213711	Endometrial stromal sarcoma				
213746	Endometrial transitional cell carcinoma	231742	Epibulbar lipodermoid-preauricular appendage-polythelia syndrome	411986	Epilepsy-cortical blindness-intellectual disability-facial dysmorphism syndrome
		83314	Epidemic typhus		
213721	Endometrial undifferentiated carcinoma	35125	Epidermal hamartoma syndrome	1946	Epilepsy-dementia-amelogenesis imperfecta syndrome
454723	Endometrioid carcinoma of ovary	35125	Epidermal nevus syndrome		
2022	Endomyocardial fibroelastosis	302	Epidermodysplasia verruciformis	1948	Epilepsy-microcephaly-skeletal dysplasia syndrome
199323	Endophthalmitis	46487	Epidermolysis bullosa acquisita		
209959	Endophthalmitis phacoanaphylactica	79404	Epidermolysis bullosa letalis	1951	Epilepsy-telangiectasia syndrome
		412181	Epidermolysis bullosa simplex due to BP230 deficiency		
2790	Endosteal hyperostosis, Worth type	412189	Epidermolysis bullosa simplex due to exophilin 5 deficiency	725	Epileptic encephalopathy with continuous spike-and-wave during slow sleep
85186	Endosteal sclerosis-cerebellar hypoplasia syndrome	158668	Epidermolysis bullosa simplex due to plakophilin deficiency	353217	Epileptic encephalopathy with global cerebral demyelination
293936	Endothelial dystrophy-iris hypoplasia-congenital cataract-stromal thinning syndrome				
		79400	Epidermolysis bullosa simplex of palms and soles	79238	Epimerase deficiency galactosemia
137602	Endotheliitis	89839	Epidermolysis bullosa simplex superficialis	1819	Epimetaphyseal skeletal dysplasia
1937	Eng-Strom syndrome	2325	Epidermolysis bullosa simplex with anodontia/hypodontia	1825	Epiphyseal dysplasia-hearing loss-dysmorphism syndrome
53540	Enhanced S-cone syndrome			1824	Epiphyseal dysplasia-microcephaly-nystagmus syndrome
60015	Enlarged parietal foramina	158681	Epidermolysis bullosa simplex with circinate migratory erythema	1952	Epiphyseal stippling syndrome-osteoclastic hyperplasia syndrome
83620	Enteric anendocrinosis				
141071	Enteric duplication cyst of the tongue	79397	Epidermolysis bullosa simplex with mottled pigmentation	399329	Epiphysiolysis of the hip
		257	Epidermolysis bullosa simplex with muscular dystrophy	399329	Epiphysiolysis of the upper femur
86880	Enteropathy-associated T-cell lymphoma	158684	Epidermolysis bullosa simplex with pyloric atresia	649	Episkopi blindness
86880	Enteropathy-type T-cell lymphoma			37612	Episodic ataxia type 1
85438	Enthesitis-related JIA	79396	Epidermolysis bullosa simplex, Dowling-Meara type	97	Episodic ataxia type 2
85438	Enthesitis-related juvenile idiopathic arthritis			79135	Episodic ataxia type 3
1177	EOCA	79396	Epidermolysis bullosa simplex, herpetiformis	79136	Episodic ataxia type 4
1177	EOCARR			211067	Episodic ataxia type 5
370334	EOE	79399	Epidermolysis bullosa simplex, Köbner type	209967	Episodic ataxia type 6
73247	EOE			209970	Episodic ataxia type 7
1020	EOFAD	79399	Epidermolysis bullosa simplex, Koebner type	401953	Episodic ataxia type 8
168829	EOPPC			37612	Episodic ataxia with myokymia
449566	Eosinophilic angiocentric fibrosis	901	Eosinophilic cellulitis	401953	Episodic ataxia with slurred speech
901	Eosinophilic cellulitis			79135	Episodic ataxia-vertigo-tinnitus-myokymia syndrome

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53583	Episodic choreoathetosis/spasticity	315	Erythrokeratoderma "en cocardes"	319	Ewing sarcoma
29822	Episodic spontaneous hypothermia	316	Erythrokeratoderma progressiva symmetrica	99734	Exercise-induced delayed-onset myotonia
93928	Epispadias	317	Erythrokeratoderma variabilis	165991	Exercise-induced hyperinsulinemic hypoglycemia
293381	Epithelial recurrent erosion dystrophy	171851	Erythrokeratoderma variabilis 3	165991	Exercise-induced hyperinsulinism
103912	Epithelio-exfoliative colitis-deafness syndrome	171851	Erythrokeratoderma variabilis, Kamouraska type	466650	Exercise-induced malignant hyperthermia
157791	Epithelioid hemangioendothelioma	317	Erythrokeratoderma variabilis, Mendes da Costa type	289586	Exfoliative ichthyosis
293202	Epithelioid sarcoma	1955	Erythrokeratoderma with ataxia	→955	Exner syndrome
254698	Epithelioid trophoblastic tumor	476096	Erythrokeratoderma-cardiomyopathy syndrome	116	Exomphalos-macroglossia-gigantism syndrome
91414	Epithelioma calcificans of Malherbe	50943	Erythrokeratolysis hiemalis	1962	Exostoses-anetoderma-brachydactyly type E syndrome
501	EPM2	318	Erythroleukemia	374	Expanded spectrum of hemifacial microsomia
263516	EPM3	1956	Erythromelalgia	322	Extrophy-epispadias complex
163696	EPM4	280379	Erythropoietic uroporphyrria associated with myeloid malignancy	440724	Extensive peripapillary myelinated nerve fibers
402082	EPM5	99977	ESCC	3294	Extensor tendons of finger anomalies
280620	EPM6	2405	Escher-Hirt syndrome	141074	External auditory canal aplasia/hypoplasia
435438	EPM7	2990	Escobar syndrome	3023	External auditory canal atresia-vertical talus-hypertelorism syndrome
424027	EPM8	2990	Escobar variant multiple pterygium syndrome	141074	External auditory canal stenosis/atresia
457265	EPM9	99976	Esophageal adenocarcinoma	231632	Extra-adrenal aldosterone-producing tumor
79278	EPP	1199	Esophageal atresia	168829	Extra-ovarian primary peritoneal carcinoma
2199	EPPK	418945	Esophageal carcinoma, salivary gland type	66662	Extracutaneous mastocytoma
→182050	Epstein syndrome	100047	Esophageal duplication cyst	182127	Extragonadal germinoma
313920	Epstein-Barr virus-associated gastric carcinoma	99977	Esophageal epidermoid carcinoma	883	Extragonadal teratoma
289661	Epstein-Barr virus-positive diffuse large B-cell lymphoma of the elderly	99977	Esophageal squamous cell carcinoma	280811	Extralobar congenital bronchopulmonary sequestration
85438	ERA	91138	Essential cryoglobulinemia	280811	Extralobar congenital pulmonary sequestration
229	Erdheim disease	2056	Essential fructosuria	2800	Extramammary Paget disease
35687	Erdheim-Chester disease	98981	Essential iris atrophy	86850	Extramedullary myeloid tumor
293381	ERED	91138	Essential mixed cryoglobulinemia	100022	Extramedullary soft tissue plasmacytoma
999	Ermine phenotype	2843	Essential pentosuria	100002	Extraneural perineurioma
160148	Eroded polypoid hyperplasia	98682	Essential strabismus	52417	Extranodal marginal zone B-cell lymphoma
→79500	Eronen-Somer-Gustafsson syndrome	3318	Essential thrombocytopenia	86879	Extranodal nasal NK/T cell lymphoma
222	Erosive pustular dermatosis of the scalp	3318	Essential thrombocytosis	370334	Extraosseous Ewing sarcoma
228264	Eruptive collagenoma	1957	Esthesioneuroblastoma	370334	Extraosseous Ewing tumor
90000	Erythema elevatum diutinum	785	Estrogen resistance syndrome	370334	Extraskeletal Ewing sarcoma
231031	Erythema palmare hereditarium	3318	ET	370334	Extraskeletal Ewing tumor
308473	Erythrocyte epimerase deficiency galactosemia	31826	Ethylene glycol poisoning	209916	Extraskeletal myxoid chondrosarcoma
308473	Erythrocyte galactose epimerase deficiency	51188	Ethylmalonic encephalopathy		
308473	Erythrocyte GALE deficiency	983	ETRS		
308473	Erythrocyte GALE-D	86880	ETTL		
171690	Erythrocyte lactate transporter defect	2892	Euhidrotic ectodermal dysplasia		
308473	Erythrocyte UDP-galactose-4-epimerase deficiency	99172	Euryblepharon		
308473	Erythrocyte uridine diphosphate galactose-4-epimerase deficiency	466682	Euthyroid Graves ophthalmopathy		
314	Erythroderma desquamativum	466682	Euthyroid Graves orbitopathy		
79394	Erythrodermic ichthyosis	1959	Evans syndrome		
247165	Erythroedema polyneuritis	444463	Evans syndrome associated with primary immunodeficiency		
		2990	EVMP5		
		251927	EVN		

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1964	Extrasystoles-short stature-hyperpigmentation-microcephaly syndrome	1778	Facial dysmorphism-shawl scrotum-joint laxity syndrome	95700	Familial adrenal hypoplasia, miniature type
		221083	Facial hemispasm	86814	Familial adult myoclonic epilepsy
251927	Extraventricular neurocytoma	3020	Facial nerve palsy due to herpes zoster infection	164736	Familial advanced sleep-phase syndrome
2725	Eye defects-arachnodactyly-cardiopathy syndrome			98880	Familial afibrinogenemia
3172	Eyebrow duplication-syndactyly syndrome	3020	Facial nerve paralysis due to VZV	1020	Familial Alzheimer disease
2985	Eyebrows and eyelashes absence-intellectual disability syndrome	85162	Facial onset sensory and motor neuronopathy	280397	Familial Alzheimer-like prion disease
		3237	Facio-audio-symphalangism	319465	Familial AML
139431	Eyelid myoclonia with and without absences	1974	Facio-digito-genital syndrome, Kuwait type	85450	Familial amyloid nephropathy
35909	F5F8D	1300	Facio-genito-popliteal syndrome	93560	Familial amyloid nephropathy due to apolipoprotein A-I variant
957	F syndrome			238269	Facio-oculo-acoustico-renal syndrome
95	FA	2048	Facio-pharyngo-glosso-masticatory diplegia		
324	Fabry disease	374	Facioauriculovertebral dysplasia	93561	Familial amyloid nephropathy due to lysozyme variant
1969	FACES syndrome				
1167	Facial asymmetry-temporal seizures syndrome	3071	Faciocutaneoskeletal syndrome	85447	Familial amyloid polyneuropathy type I (Portuguese-Swedish-Japanese Type)
141051	Facial dermoid cyst	915	Faciodigitogenital syndrome		
480701	Facial diplegia with paresthesias	915	Faciogenital dysplasia	85448	Familial amyloid polyneuropathy type IV
480701	Facial diplegia with paresthesias variant of GBS	269	Facioscapulohumeral dystrophy		
480701	Facial diplegia with paresthesias variant of Guillain-Barré syndrome	269	Facioscapulohumeral muscular dystrophy	85448	Familial amyloidosis, Finnish type
				269	Facioscapulohumeral myopathy
→3157	Facial dysmorphism-ambiguous genitalia-hypopituitarism-short limbs syndrome	98879	Factor IX deficiency	199279	Familial angiolipomatosis
		220436	Factor V Quebec	91378	Familial angioneurotic edema
1969	Facial dysmorphism-anorexia-cachexia-eye and skin anomalies syndrome	98878	Factor VIII deficiency	229	Familial aortic dissection
		300359	FACU	425	Familial apoA-I deficiency
284169	Facial dysmorphism-developmental delay-behavioral abnormalities syndrome due to 10p11.21p12.31 microdeletion	306550	FADD-related immunodeficiency	309020	Familial apoC-II deficiency
		994	FADS	309020	Familial apolipoprotein C-II deficiency
466950	Facial dysmorphism-developmental delay-behavioral abnormalities syndrome due to WAC point mutation	882	FAH deficiency	1416	Familial articular chondrocalcinosis
		329308	FAHN	334	Familial atrial fibrillation
352712	Facial dysmorphism-immunodeficiency-livedo-short stature syndrome	→168569	Faisalabad histiocytosis	615	Familial atrial myxoma
		3304	Fallot complex-intellectual disability-growth delay syndrome	436242	Familial atrial tachyarrhythmia-infra-Hisian cardiac conduction disease
86814	FAME	300359	Familial atypical cold urticaria		
2588	Facial dysmorphism-intellectual disability-short stature-hearing loss syndrome	397685	Familial hyperprolactinemia	404560	Familial atypical mole syndrome
		86	Familial abdominal aortic aneurysm	404560	Familial atypical multiple mole melanoma syndrome
412022	Facial dysmorphism-lens dislocation-anterior segment abnormalities-nontraumatic conjunctive cysts syndrome	88619	Familial acute necrotizing encephalopathy	404560	Familial atypical multiple mole melanoma-pancreatic carcinoma syndrome
		733	Familial adenomatous polyposis		
412022	Facial dysmorphism-lens dislocation-anterior segment abnormalities-spontaneous filtering blebs syndrome	261584	Familial adenomatous polyposis due to 5q22.2 microdeletion	86820	Familial avascular necrosis of femoral head
		261584	Familial adenomatous polyposis due to del(5)(q22.2)	2398	Familial benign cervical lipomatosis
1970	Facial dysmorphism-macrocephaly-myopia-Dandy-Walker malformation syndrome	261584	Familial adenomatous polyposis due to monosomy 5q22.2	2841	Familial benign chronic pemphigus
		404	Familial adrenal adenoma	1551	Familial benign copper deficiency
1970	Facial dysmorphism-macrocephaly-myopia-Dandy-Walker malformation syndrome	95700	Familial adrenal hypoplasia with absent pituitary LH	363989	Familial benign flecked retina
		95700	Familial adrenal hypoplasia with absent pituitary luteinizing hormone	405	Familial benign hypercalcemia
				405	Familial benign hypocalciuric hypercalcemia

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
231160	Familial berry aneurysm	86814	Familial cortical myoclonic tremor and epilepsy	464756	Familial gastric type 1 neuroendocrine tumor
402075	Familial bicuspid aortic valve			231040	Familial generalized lentiginosis
221061	Familial brain cavernous angioma	319189	Familial cortical myoclonus	99819	Familial gestational hyperthyroidism
221061	Familial brain cavernous hemangioma	1416	Familial CPPD		
		85453	Familial cutaneous amyloidosis		
227535	Familial breast cancer	53296	Familial cutaneous collagenoma	361	Familial glucocorticoid deficiency
227535	Familial breast carcinoma	313846	Familial cutaneous telangiectasia and oropharyngeal predisposition cancer syndrome	3000	Familial gonadotropin-independent male-limited sexual precocity
36382	Familial CAD			540	Familial hemophagocytic lymphohistiocytosis
2678	Familial café-au-lait spots				
1416	Familial calcium pyrophosphate deposition	211	Familial cylindromatosis	32960	Familial Hibernian fever
		97345	Familial dementia, British type		
1768	Familial caudal dysgenesis	97346	Familial dementia, Danish type	540	Familial HLH
464760	Familial cavitory optic disc anomaly	313808	Familial dementia, Neumann type	2604	Familial hollow visceral myopathy
1416	Familial CC	1799	Familial developmental dysphasia	403	Familial hyperaldosteronism type 1
169085	Familial CD8 deficiency	26106	Familial diffuse cancer of stomach	404	Familial hyperaldosteronism type 2
892	Familial cerebelloretinal angiomatosis	26106	Familial diffuse gastric cancer	251274	Familial hyperaldosteronism type 3
		85169	Familial digital arthropathy-brachydactyly	403	Familial hyperaldosteronism type I
439254	Familial cerebral amyloid angiopathy			300751	Familial dilated cardiomyopathy with conduction defect due to LMNA mutation
		251274	Familial hyperaldosteronism type III		
221061	Familial cerebral cavernoma			79506	Familial hyperalphalipoproteinemia
221061	Familial cerebral cavernous malformation	18	Familial distal primary acidosis	94086	Familial hypercalcemia-nephrocalcinosis-indicanuria syndrome
231160	Familial cerebral saccular aneurysm	85192	Familial doughnut lesions of skull		
36382	Familial cervical artery dissection	75376	Familial drusen	238475	Familial hypercholanemia
481662	Familial Chilblain lupus	79142	Familial Dupuytren contracture	178345	Familial hyperestrogenism
1428	Familial chondromalacia patellae	1764	Familial dysautonomia	757	Familial hyperkalemic hypertension
444490	Familial chylomicronemia syndrome	314381	Familial dysautonomia with contractures	682	Familial hyperkalemic periodic paralysis
404560	Familial Clark nevus syndrome				
293144	Familial clubfoot due to 5q31 microdeletion	412	Familial dysbetalipoproteinemia	412	Familial hyperlipoproteinemia type 3
		98881	Familial dysfibrinogenemia		
238578	Familial clubfoot due to 17q23.1q23.2 microduplication	324588	Familial dyskinesia and facial myokymia	306661	Familial hyperphosphatemic tumoral calcinosis/Hyperphosphatemic hyperostosis syndrome
293150	Familial clubfoot due to PITX1 point mutation	404560	Familial dysplastic nevus syndrome		
199315	Familial clubfoot with or without associated lower limb anomalies	1885	Familial ectopia lentis	682	Familial hyperPP
		2762	Familial ectopic ossification		
464760	Familial CODA	85110	Familial encephalopathy with neuroserpin inclusion bodies	99763	Familial hyperreninemic hypoadosteronism type 1
47045	Familial cold autoinflammatory syndrome	101039	Familial epilepsy and mental retardation limited to females	99764	Familial hyperreninemic hypoadosteronism type 2
247868	Familial cold autoinflammatory syndrome type 2	391384	Familial episodic pain syndrome	424	Familial hyperthyroidism due to mutations in TSH receptor
47045	Familial cold urticaria	391392	Familial episodic pain syndrome with predominantly lower limb involvement	427	Familial hypoadosteronism
300359	Familial cold urticaria with common variable immunodeficiency			425	Familial hypoalphalipoproteinemia
440437	Familial colorectal cancer Type X	391389	Familial episodic pain syndrome with predominantly upper body involvement	405	Familial hypocalciuric hypercalcemia
238722	Familial congenital controlateral synkinesia	90042	Familial erythrocytosis	93372	Familial hypocalciuric hypercalcemia type 1
95494	Familial congenital hypopituitarism	225968	Familial essential thrombocythemia	101049	Familial hypocalciuric hypercalcemia type 2
238722	Familial congenital mirror movements	85195	Familial expansile osteolysis		
		891	Familial exudative vitreoretinopathy	101050	Familial hypocalciuric hypercalcemia type 3
451612	Familial congenital nasolacrimal duct obstruction	98820	Familial focal epilepsy with variable foci	248408	Familial hypodysfibrinogenemia
91498	Familial congenital palsy of trochlear nerve	314022	Familial fundic gland polyposis with gastric cancer	101041	Familial hypofibrinogenemia
				440	Familial hypospadias
				225154	Familial IBSN

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
1677	Familial idiopathic dilatation of the right atrium	293910	Familial isolated arrhythmogenic ventricular cardiomyopathy, classic form	217330	Familial juvenile hyperuricemic nephropathy type 2
656	Familial idiopathic nephrotic syndrome			493	Familial keratoacanthoma
656	Familial idiopathic steroid-resistant nephrotic syndrome	293888	Familial isolated arrhythmogenic ventricular cardiomyopathy, left dominant form	293936	Familial keratoconus with cataract
93214	Familial idiopathic steroid-resistant nephrotic syndrome with diffuse mesangial proliferation			3267	Familial lambdoid synostosis
		79293	Familial LCAT deficiency		
93217	Familial idiopathic steroid-resistant nephrotic syndrome with diffuse mesangial sclerosis	293910	Familial isolated arrhythmogenic ventricular cardiomyopathy, right dominant form	523	Familial leiomyomatosis and renal cell cancer
				523	Familial leiomyomatosis cutis et uteri
93213	Familial idiopathic steroid-resistant nephrotic syndrome with focal segmental glomerulosclerosis	217656	Familial isolated arrhythmogenic ventricular dysplasia	523	Familial leiomyomatosis with renal carcinoma
				293899	Familial isolated arrhythmogenic ventricular dysplasia, biventricular form
93213	Familial idiopathic steroid-resistant nephrotic syndrome with focal segmental hyalinosis	293910	Familial isolated arrhythmogenic ventricular dysplasia, classic form	871	Familial Lenègre disease
				293888	Familial isolated arrhythmogenic ventricular dysplasia, left dominant form
93216	Familial idiopathic steroid-resistant nephrotic syndrome with minimal changes	293910	Familial isolated arrhythmogenic ventricular dysplasia, right dominant form	871	Familial Lev-Lenègre disease
				309015	Familial lipoprotein lipase deficiency
225154	Familial infantile bilateral striatal necrosis	217656	Familial isolated ARVC	768	Familial long QT syndrome
300373	Familial infantile gigantism	217656	Familial isolated ARVD	75381	Familial macular edema
448348	Familial infantile gigantism due to a point mutation	295014	Familial isolated clinodactyly of fingers	3000	Familial male-limited precocious puberty
448372	Familial infantile gigantism due to dup(X)q(26)	101351	Familial isolated congenital asplenia	401942	Familial median cleft of the upper and lower lips
448372	Familial infantile gigantism due to Xq26 microduplication	154	Familial isolated dilated cardiomyopathy	342	Familial Mediterranean fever
300547	Familial infantile hypercalcemia with suppressed intact parathyroid hormone	99879	Familial isolated hyperparathyroidism	99361	Familial medullary thyroid carcinoma
		2238	Familial isolated hypoparathyroidism	35858	Familial megaloblastic anemia
352582	Familial infantile myoclonic epilepsy	2239	Familial isolated hypoparathyroidism due to agenesis of parathyroid gland	618	Familial melanoma
352582	Familial infantile myoclonus epilepsy			165805	Familial mesial temporal lobe epilepsy with febrile seizures
225154	Familial infantile striatonigral degeneration	189466	Familial isolated hypoparathyroidism due to impaired PTH secretion	741	Familial mitral valve prolapse
225154	Familial infantile striatonigral necrosis	314777	Familial isolated pituitary adenoma	276399	Familial MNG
2454	Familial intestinal malrotation-facial anomalies syndrome	397685	Familial isolated prolactin receptor deficiency	99361	Familial MTC
2300	Familial intestinal polyatresia syndrome	75249	Familial isolated restrictive cardiomyopathy	276399	Familial multinodular goiter
231160	Familial intracranial saccular aneurysm	411788	Familial isolated trichomegaly	35909	Familial multiple coagulation factor deficiency
		96	Familial isolated vitamin E deficiency	523	Familial multiple cutaneous leiomyomas
217656	Familial isolated arrhythmogenic right ventricular cardiomyopathy	2295	Familial joint instability syndrome	338	Familial multiple fibrofolliculoma
217656	Familial isolated arrhythmogenic right ventricular dysplasia	2295	Familial joint laxity	500	Familial multiple lentiginos syndrome
217656	Familial isolated arrhythmogenic ventricular cardiomyopathy	180176	Familial juvenile gigantomastia	231040	Familial multiple lentiginos syndrome without systemic involvement
		209886	Familial juvenile gouty nephropathy		
293899	Familial isolated arrhythmogenic ventricular cardiomyopathy, biventricular form	180176	Familial juvenile hypertrophy of the breast	199276	Familial multiple lipomatosis
		209886	Familial juvenile hyperuricemic nephropathy type 1	263662	Familial multiple meningioma
				624	Familial multiple nevi flammei
				624	Familial multiple port-wine stains
				867	Familial multiple trichoepithelioma
				922	Familial nasal acilia
				209886	Familial nephropathy with gout
				424	Familial non-immune hyperthyroidism

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306658	Familial normophosphatemic tumoral calcinosis	71290	Familial platelet syndrome with predisposition to acute myelogenous leukemia	93560	Familial renal amyloidosis due to apolipoprotein A-I variant
280403	Familial omphalocele syndrome with facial dysmorphism	330061	Familial polymorphous light eruption of American Indians	238269	Familial renal amyloidosis due to apolipoprotein A-II variant
154	Familial or idiopathic dilated cardiomyopathy	733	Familial polyposis coli	93561	Familial renal amyloidosis due to lysozyme variant
75249	Familial or idiopathic restrictive cardiomyopathy	261584	Familial polyposis coli due to monosomy 5q22.2	69076	Familial renal glucosuria
569	Familial or sporadic hemiplegic migraine	99810	Familial porencephaly	284247	Familial retinal arterial macroaneurysm
		443062	Familial porphyria cutanea tarda	231108	Familial rhabdoid tumor
443236	Familial orthostatic tachycardia due to norepinephrine transporter deficiency	2196	Familial primary hypomagnesemia with hypercalciuria and nephrocalcinosis with severe ocular involvement	→168569	Familial Rosaï-Dorfman disease
435329	Familial ossifying fibroma			168624	Familial scaphocephaly syndrome, McGillivray type
251262	Familial osteochondritis dissecans	31043	Familial primary hypomagnesemia with hypercalciuria and nephrocalcinosis without severe ocular involvement	171839	Familial scaphocephaly-radioulnar synostosis syndrome
2769	Familial osteodysplasia, Anderson type			3135	Familial Scheuermann disease
2801	Familial osteoectasia			3135	Familial Scheuermann juvenile kyphosis
86820	Familial osteonecrosis of the femoral head	34527	Familial primary hypomagnesemia with normocalciuria and normocalcemia	481986	Familial schizencephaly
79093	Familial osteosclerosis with abnormalities of the nervous system and meninges	353220	Familial primary localized cutaneous amyloidosis	→168569	Familial SHML
1333	Familial pancreatic cancer	2257	Familial primary pulmonary hypoplasia	51083	Familial short QT syndrome
1333	Familial pancreatic carcinoma			166282	Familial sick sinus syndrome
319487	Familial papillary or follicular thyroid carcinoma	65748	Familial primary self-healing squamous epithelioma of the skin, Ferguson-Smith type	→168569	Familial sinus histiocytosis with massive lymphadenopathy
97290	Familial papillary thyroid carcinoma with renal papillary neoplasia	871	Familial progressive cardiac conduction defect	166282	Familial sinus node dysfunction
99877	Familial parathyroid adenoma	871	Familial progressive heart block	300345	Familial SLE
99878	Familial parathyroids hyperplasia			3135	Familial spinal osteochondrosis
97	Familial paroxysmal ataxia	280628	Familial progressive hyper- and hypopigmentation	2903	Familial spontaneous pneumothorax
98809	Familial paroxysmal kinesigenic dyskinesia	79146	Familial progressive hyperpigmentation	3197	Familial startle disease
342	Familial paroxysmal polyserositis	313808	Familial progressive subcortical gliosis		Familial steroid-resistant nephrotic syndrome with sensorineural deafness
228140	Familial paroxysmal ventricular fibrillation, non Brugada type	1767	Familial progressive vestibulocochlear dysfunction	1325	Familial streblodactyly with amino-aciduria
98820	Familial partial epilepsy with variable foci	1331	Familial prostate cancer	2456	Familial supernumerary nipples
79084	Familial partial lipodystrophy type 1	90044	Familial pseudohyperkalemia	370034	Familial syringomyelia
2348	Familial partial lipodystrophy type 2	→3202	Familial pseudohyperkalemia type 1	300345	Familial systemic lupus erythematosus
79083	Familial partial lipodystrophy type 3			91387	Familial TAAD
2348	Familial partial lipodystrophy, Dunnigan type	2989	Familial pterygium of the conjunctiva	98819	Familial temporal epilepsy
79084	Familial partial lipodystrophy, Köbberling type	275777	Familial pulmonary arterial hypertension	91387	Familial thoracic aortic aneurysm and aortic dissection
466729	Familial patent arterial duct	319487	Familial pure nonmedullary thyroid carcinoma	71493	Familial thrombocytopenia
871	Familial PCCD	1675	Familial pyrimidinemia	71493	Familial thrombocytosis
93333	Familial pelvis-scapular dysplasia	79147	Familial reactive perforating collagenosis	329319	Familial thrombocytosis with transverse limb defect
29072	Familial pheochromocytoma-paraganglioma	46348	Familial rectal pain	3324	Familial thrombomodulin anomalies
98809	Familial PKD	69126	Familial recurrent arthritis	93953	Familial thyroglossal duct cyst
		2809	Familial recurrent peripheral facial palsy	95716	Familial thyroid dysmorphogenesis
71290	Familial platelet disorder with associated myeloid malignancy	85450	Familial renal amyloidosis	53372	Familial trembling of the chin
				93583	Familial TTP
				53715	Familial tumoral calcinosis
				289365	Familial vesicoureteral reflux
				2604	Familial visceral myopathy

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2808	Familial vocal cord dysfunction			→182050	Fechtner syndrome
289365	Familial VUR	289527	Fatal infantile hypertrophic cardiomyopathy due to NADH-CoQ reductase deficiency	79292	FED
170	Familial woolly hair syndrome			247165	Feer disease
170	Familial woolly hair syndrome	17	Fatal infantile lactic acidosis with methylmalonic aciduria	98969	Fehr corneal dystrophy
404560	FAMM-PC syndrome			1192	Feigenbaum-Bergeron-Richardson syndrome
404560	FAMMM syndrome			1305	Feingold syndrome
84	Fanconi anemia	168566	Fatal mitochondrial disease due to combined oxidative phosphorylation defect type 3	391641	Feingold syndrome type 1
84	Fanconi pancytopenia			391646	Feingold syndrome type 2
→2697	Fanconi syndrome-ichthyosis-dysmorphism syndrome	168566	Fatal mitochondrial disease due to COXPD3	53693	Fellman disease
2088	Fanconi-Bickel disease	401869	Fatal multiple mitochondrial dysfunctions syndrome type 1	47612	Felty syndrome
733	FAP			404466	Female infertility due to zona pellucida defect
261584	FAP due to monosomy 5q22.2	401874	Fatal multiple mitochondrial dysfunctions syndrome type 2	2973	Female pseudohermaphroditism-anorectal anomalies syndrome
2792	Fara-Chlupackova syndrome	363424	Fatal multiple mitochondrial dysfunctions syndrome type 3	2975	Female pseudohermaphroditism-skeletal anomalies syndrome
333	Farber disease			101039	Female restricted epilepsy with intellectual disability
333	Farber lipogranulomatosis	457406	Fatal multiple mitochondrial dysfunctions syndrome type 4	1987	Femoral agenesis/hypoplasia
99906	Farmer's lung disease			295067	Femoral agenesis/hypoplasia, bilateral
1915	FAS	391343	Fatal post-viral neurodegenerative disorder	295065	Femoral agenesis/hypoplasia, unilateral
3261	FAS deficiency	480773	FATCO syndrome	399329	Femoral head epiphysiolysis
1915	FASD	816	Fatty acid alcohol oxidoreductase deficiency	1988	Femoral hypoplasia-unusual facies syndrome
164736	FASPS			1987	Femoral intercalary meromelia
166105	FASTKD2-related infantile mitochondrial encephalomyopathy	329308	Fatty acid hydroxylase-associated neurodegeneration	295067	Femoral intercalary meromelia, bilateral
439854	Fatal congenital hypertrophic cardiomyopathy due to glycogen storage disease	2064	Faulk-Epstein-Jones syndrome	295065	Femoral intercalary meromelia, unilateral
439854	Fatal congenital hypertrophic cardiomyopathy due to glycogenosis	→97229	Fazio-Londe disease	1988	Femoral intercalary meromelia, bilateral
439854	Fatal congenital hypertrophic cardiomyopathy due to GSD	405	FBH	295067	Femoral intercalary meromelia, bilateral
466	Fatal familial insomnia	405	FBHH	295065	Femoral intercalary meromelia, unilateral
1561	Fatal infantile cardioencephalomyopathy due to cytochrome C oxidase deficiency	404451	FBLN1-related developmental delay-central nervous system anomaly-syndactyly syndrome	1988	Femoral-facial syndrome
1561	Fatal infantile COX deficiency	47045	FCAS	294977	Femrotibiofibular intercalary transverse meromelia
1561	Fatal infantile cytochrome C oxidase deficiency	247868	FCAS2	295091	Femrotibiofibular intercalary transverse meromelia, bilateral
166073	Fatal infantile encephalopathy with mitochondrial respiratory chain defects	440437	FCCTX	295089	Femrotibiofibular intercalary transverse meromelia, unilateral
166063	Fatal infantile encephalopathy with olivopontocerebellar hypoplasia	98970	FCD	2019	Femur-fibula-ulna complex
→370114	Fatal infantile encephalopathy-pulmonary hypertension syndrome	268961	FCD type I	2019	Femur-fibula-ulna dysostosis
289527	Fatal infantile HCM due to mitochondrial complex I deficiency	268973	FCD type Ia	2019	Femur-fibula-ulna syndrome
280553	Fatal infantile hypertonic myofibrillar myopathy	268980	FCD type Ib	60015	Fenestrae parietales symmetricae
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	268987	FCD type Ic	85110	FENIB
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	268994	FCD type II	1184	Fenton-Wilkinson-Toselano syndrome
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	269001	FCD type IIa	45358	FEOM
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	269008	FCD type IIb	391384	FEPS
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	272	FCMD	65748	Ferguson-Smith disease
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	86814	FCMTE	2180	Ferlini-Ragno-Calzolari syndrome
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	99654	FCPD	157846	Ferritin-related neurodegeneration
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	3071	FCS syndrome	397922	Ferro-cerebro-cutaneous syndrome
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	47045	FCU	139491	Ferroportin disease
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	324	FD	40366	Fetal acitretin/etretinate syndrome
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	324588	DFDM		
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	26106	FDGC		
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	412022	FDLAB syndrome		
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	163703	Febrile infection-related epilepsy syndrome		
289527	Fatal infantile hypertrophic cardiomyopathy due to mitochondrial complex I deficiency	98974	FECD		

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994	Fetal akinesia deformation sequence	1988	FFS	480773	Fibular aplasia-tibial campomelia-oligosyndactyly syndrome
		2019	FFU complex		
363409	Fetal akinesia-cerebral and retinal hemorrhage syndrome	93932	FG syndrome type 1	1757	Fibular dimelia-diplopodia syndrome
		313855	FGFR2-related bent bone dysplasia		
1915	Fetal alcohol spectrum disorders	1305	FGLDS	93323	Fibular hemimelia
1915	Fetal alcohol syndrome	391641	FGLDS1	295083	Fibular hemimelia, bilateral
1908	Fetal aminopterin syndrome	391646	FGLDS2	295081	Fibular hemimelia, unilateral
1041	Fetal anasarca	403	FH1	2854	Fibular hypoplasia or aplasia-femoral bowing-oligodactyly syndrome
853	Fetal and neonatal alloimmune thrombocytopenia	404	FH2		
		251274	FH3		
370076	Fetal carbamazepine syndrome	403	FH-I	93323	Fibular longitudinal meromelia
1911	Fetal cocaine syndrome	404	FH-II	295083	Fibular longitudinal meromelia, bilateral
294	Fetal cytomegalovirus syndrome	251274	FH-III		
1912	Fetal dihydantoin syndrome	→168569	FHC	295081	Fibular longitudinal meromelia, unilateral
465824	Fetal encasement syndrome	401920	FHCC		
97360	Fetal face syndrome	405	FHH	2256	Fibulo-ulnar hypoplasia-renal anomalies syndrome
85212	Fetal Gaucher disease	93372	FHH type 1		
1912	Fetal hydantoin syndrome	101049	FHH type 2	79306	FC1 deficiency
1041	Fetal hydrops	101050	FHH type 3	29207	Fiessinger-Leroy disease
1909	Fetal indomethacin syndrome	99763	FHHA1	29207	Fiessinger-Leroy-Reiter syndrome
1910	Fetal iodine syndrome	99764	FHHA2	2756	Figuera syndrome
1055	Fetal left ventricular aneurysm	2196	FHHNC with severe ocular involvement	99879	FIHPT
284362	Fetal lung interstitial tumor				
1917	Fetal methylmercury syndrome	31043	FHHNC without severe ocular involvement	3255	Filippi syndrome
1918	Fetal minoxidil syndrome				
295	Fetal parvovirus syndrome	263479	FHI	352712	FILS syndrome
290	Fetal rubella syndrome	397618	FHONDA syndrome	352582	FIME
3312	Fetal thalidomide syndrome	1988	FHUF5	1272	Fine-Lubinsky syndrome
1913	Fetal trimethadione syndrome	251601	Fibrillary astrocytoma	369979	Finger hyperphalangy-toe anomalies-severe pectus excavatum syndrome
1906	Fetal valproate syndrome	331	Fibrin-stabilizing factor deficiency	97232	Fingerprint body myopathy
1906	Fetal valproic acid syndrome	93562	Fibrinogen A alpha-chain amyloidosis	209335	Finkel disease
1914	Fetal warfarin syndrome				
166068	Fetal-onset olivopontocerebellar hypoplasia	439881	Fibrinous bronchitis	2036	Finlay-Marks syndrome
		477650	Fibroblastic rheumatism	839	Finnish congenital nephrosis
95431	Feto-fetal transfusion syndrome	99654	Fibrocalculous pancreatic diabetes	609	Finnish tibial muscular dystrophy
69063	Fetomaternal alloimmunization with antenatal glomerulopathies	99654	Fibrocalculous pancreatopathy	399086	Finnish upper limb-onset distal myopathy
		2021	Fibrochondrogenesis	1825	Finucane-Kurtz-Scott syndrome
464724	Fever-associated acute infantile liver failure syndrome	337	Fibrodysplasia ossificans progressiva	314777	FIPA
				163703	FIRES
163703	Fever-induced refractory epileptic encephalopathy in school-aged children	122	Fibrolamellar hepatocarcinoma	141136	First branchial arch syndrome
				401920	Fibrolamellar hepatocellular carcinoma
891	FEVR	401920	Fibrolamellar hepatocellular carcinoma	141013	First branchial cleft anomaly
254492	FFA				
398166	FFDD	79105	Fibromyxosarcoma	141013	First branchial cleft cyst
79133	FFDD1	84090	Fibronectin glomerulopathy	141013	First branchial cleft fistula
398173	FFDD2	2030	Fibrosarcoma	79292	Fish-eye disease
1807	FFDD3	63999	Fibrosing mediastinitis	98919	Fisher syndrome
398189	FFDD4	249	Fibrous dysplasia of bone	840	Fistulous vegetative verrucous hydradenoma
79133	FFDD type I				
398173	FFDD type II	2639	Fibular aplasia-complex brachydactyly syndrome	2823	Fitzsimmons-Guilbert syndrome
1807	FFDD type III				
398189	FFDD type IV	1118	Fibular aplasia-ectrodactyly syndrome	2824	Fitzsimmons-McLachlan-Gilbert syndrome
98820	FFEVF				
				2820	Fitzsimmons-Walson-Mellor syndrome
				293812	Fixed pigmented erythema
				3092	Fixed subaortic stenosis
				209886	FJHN type 1

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
217330	FJHN type 2	443804	Focal stiff-person syndrome	353220	FPLCA
1968	Flat face-microstomia-ear anomaly syndrome	79093	Foix-Alajouanine syndrome	79084	FPLD1
		2048	Foix-Chavany-Marie syndrome	2348	FPLD2
79293	FLD	79097	Folinic acid-responsive seizures	79083	FPLD3
98970	Fleck corneal dystrophy	113	Follicular atrophoderma and basal cell carcinomas	280356	FPLD4
409	Flegel disease			435651	FPLD5
284362	FLIT	79459	Follicular atrophoderma-basal cell carcinoma	435660	FPLD6
2044	Floating-Harbor syndrome			71290	FPS/AML syndrome
83451	Florid cemento-osseous dysplasia	300552	Follicular cholangitis and pancreatitis	313808	FPSG
83451	Florid osseous dysplasia			69126	FRA
2045	FLOTCH syndrome	86902	Follicular dendritic cell sarcoma	908	Fragile X syndrome
199306	FLP	69745	Follicular dyskeratoma	93256	Fragile X-associated tremor/ataxia syndrome
99734	Fluctuating myotonia	2112	Follicular hamartoma-alopecia-cystic fibrosis syndrome		
1685	Fluke infection			525	Follicular lichen planus
2047	Flynn-Aird syndrome	545	Follicular lymphoma	2523	Franek-Bocker-Kahlen syndrome
69063	FMAIG	300552	Follicular pancreatocholangitis	137834	Frank-Ter Haar syndrome
342	FMF			100026	Franklin disease
276399	FMNG	243	Follicular stimulating hormone-resistant ovaries	2108	François dyscephalic syndrome
3000	FMPP	79100	Folliculitis ulerythematosia reticulata	79149	François syndrome
319487	FNMTC			178512	Folliculotropic mycosis fungoides
137675	Foamy myocardial transformation of infancy	228371	Foodborne botulism		
2143	FOAR syndrome	3454	Foot contractures-muscle atrophy-oculomotor apraxia syndrome	→2052	Fraser-like syndrome
308013	Focal acral hyperkeratosis			337	FOP
83451	Focal cemento-osseous dysplasia	60015	Foramina parietalia permagna	908	FraX syndrome
2092	Focal dermal hypoplasia	366	Forbes disease	908	FRAXA syndrome
352587	Focal epilepsy-intellectual disability-cerebro-cerebellar malformation	141071	Foregut duplication cyst of the tongue	100973	FRAXE intellectual disability
352587	Focal epilepsy-intellectual disability-dysarthria-ataxia syndrome			100974	FRAXF syndrome
398166	Focal facial dermal dysplasia	51208	Formiminoglutamic aciduria	95	FRDA
79133	Focal facial dermal dysplasia 1, Brauer type	51208	Formiminotransferase cyclodeaminase deficiency	834	Free sialic acid storage disease
		3238	Forney syndrome	309324	Free sialic acid storage disease, infantile form
398173	Focal facial dermal dysplasia 2, Brauer-Setleis type	3238	Forney-Robinson-Pascoe syndrome	2053	Freeman-Sheldon syndrome
		178333	Forsius-Eriksson syndrome	1147	Freeman-Sheldon syndrome variant
1807	Focal facial dermal dysplasia 3, Setleis type	178333	Forsius-Eriksson type ocular albinism	2673	Freire Maia-Pinheiro-Opitz syndrome
		85162	FOSMN syndrome	2723	Freire-Maia syndrome
398189	Focal facial dermal dysplasia 4	3219	Fountain syndrome	→264200	Frias syndrome
79133	Focal facial dermal dysplasia type I	141037	Fourth branchial cleft anomaly	85335	Fried syndrome
398173	Focal facial dermal dysplasia type II	141037	Fourth branchial cleft cyst	2487	Fried-Goldberg-Mundel syndrome
1807	Focal facial dermal dysplasia type III	141037	Fourth branchial cleft fistula	1969	Friedman-Goodman syndrome
398189	Focal facial dermal dysplasia type IV	397618	Foveal hypoplasia-optic nerve decussation defect-anterior segment dysgenesis syndrome	95	Friedreich ataxia
398189	Focal facial preauricular dysplasia			96	Friedreich-like ataxia
221083	Focal myoclonus of face	2253	Foveal hypoplasia-presenile cataract syndrome	99672	Fried's tooth and nail syndrome
48918	Focal myositis			1931	Frontal encephalocele
48918	Focal nodular myositis	221126	Fowler syndrome	254492	Frontal fibrosing alopecia
2200	Focal palmoplantar and gingival hyperkeratosis	2795	Fowler-Christmas-Chapple syndrome	1791	Frontofacionasal dysplasia
				1799	FOXP2-associated dysphasia
370002	Focal palmoplantar keratoderma with joint keratoses	275777	FPAH	141168	Frontonasal arteriovenous malformation
		71290	FPD/AML syndrome		
443804	Focal stiff limb syndrome	280628	FPHH		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
228390	Frontonasal dysplasia with alopecia and genital anomaly	622	Functional methionine synthase deficiency	79237	GALK deficiency
				79237	GALK-D
306542	Frontonasal dysplasia-severe microphthalmia-severe facial clefting syndrome	308380	Functional methionine synthase deficiency type cbIDv1	100086	Gallbladder neuroendocrine tumor
		2169	Functional methionine synthase deficiency type cbIE	2065	Galloway syndrome
391474	Frontorhiny			2065	Galloway-Mowat syndrome
275872	Frontotemporal dementia with amyotrophic lateral sclerosis	2170	Functional methionine synthase deficiency type cbIG	309297	GALNS deficiency
				79239	GALT deficiency
275872	Frontotemporal dementia with motor neuron disease	91348	Functioning gonadotropic adenoma	2325	Gamborg-Nielsen syndrome
		91348	Functioning pituitary gonadotropic adenoma	3035	Game-Friedman-Paradise syndrome
293848	Frontotemporal dementia, right temporal atrophy variant	227796	Fundus albipunctatus	2066	Gamma-aminobutyric acid transaminase deficiency
2141	Froster-Huch syndrome	827	Fundus flavimaculatus	212	Gamma-cystathionase deficiency
2215	Froster-Iskenius-Watson-Hall syndrome	99004	Fundus pulverulentus	33573	Gamma-glutamyl transpeptidase deficiency
		207000	Fungal myositis		
2056	Fructokinase deficiency	→60030	Furlong syndrome	33574	Gamma-glutamylcysteine synthetase deficiency
348	Fructose-1,6-bisphosphatase deficiency	2579	Furukawa-Takagi-Nakao syndrome	100026	Gamma-HCD
		591	Furuncular myiasis	100026	Gamma-heavy chain disease
2057	Frydman-Cohen-Karmon syndrome	591	Furunculoid myiasis	22	Gamma-hydroxybutyric aciduria
2429	Fryns macrocephaly	591	Furunculoid myiasis	353	Gamma-sarcoglycanopathy
1104	Fryns microphthalmia syndrome	228119	Fusariosis	682	Gamstorp disease
2059	Fryns syndrome	228119	Fusarium infection	682	Gamstorp episodic adynamy
→2995	Fryns-Aftimos syndrome	2287	Fused mandibular incisors	382	GAMT deficiency
2497	Fryns-Hofkens-Fabry syndrome	2498	Fusion of metacarpals 4 and 5	643	GAN
2058	Fryns-Smeets-Thiry syndrome	35909	FV and FVIII combined deficiency	251937	Gangliocytoma
1305	FS	908	FXS	251949	Ganglioglioma
391641	FS1	93256	FXTAS syndrome	251877	Ganglioneuroblastoma
391646	FS2	364	G6P deficiency	251992	Ganglioneuroma
269	FSH dystrophy	79258	G6P deficiency type 1a	2067	GAPO syndrome
243	FSH-RO	79259	G6P deficiency type 1b	314022	GAPPS
269	FSHD	79259	G6P translocase deficiency	3469	Garcia-Lurie syndrome
51208	FTCD deficiency	79259	G6PT deficiency	79665	Gardner syndrome
275872	FTD-ALS	25	GA1	324636	Gardner-Diamond syndrome
275872	FTD-MND	2066	GABA transaminase deficiency	2075	Gardner-Silengo-Wachtel syndrome
247790	FTH1-associated iron overload	79402	GABEB	99000	Gass disease
247790	FTH1-related iron overload	90041	Gaisböck syndrome		
98974	Fuchs endothelial corneal dystrophy	487	Galactocerebrosidase deficiency	314022	Gastric adenocarcinoma and proximal polyposis of the stomach
263479	Fuchs heterochromic iridocyclitis	79237	Galactokinase deficiency	423781	Gastric carcinoma, salivary gland type
349	Fucosidosis	79237	Galactokinase deficiency galactosemia	141071	Gastric duplication cyst of the tongue
2854	Fuhrmann syndrome			332	Gastric intrinsic factor deficiency
2854	Fuhrmann-Rieger-de Sousa syndrome	309297	Galactosamine-6-sulfatase deficiency	36273	Gastric linitis plastica
→85199	Fukuda-Miyanomae-Nakata syndrome	79238	Galactose epimerase deficiency	100075	Gastric neuroendocrine tumor
551	Fukuhara syndrome	79239	Galactose-1-phosphate uridylyltransferase deficiency	481469	Gastric neuroendocrine tumor type 1
272	Fukuyama congenital muscular dystrophy	79239	Galactosemia type 1	481475	Gastric neuroendocrine tumor type 2
		79237	Galactosemia type 2		
90062	Fulminant hepatic failure	79238	Galactosemia type 3	481478	Gastric neuroendocrine tumor type 3
35063	Fulminant viral hepatitis	351	Galactosialidosis		
24	Fumarase deficiency	487	Galactosylceramidase deficiency	481481	Gastric neuroendocrine tumor type 4
24	Fumaric aciduria	75496	Galactosyltransferase I deficiency		
882	Fumarylacetoacetase deficiency	487	GALC deficiency	418959	Gastric squamous cell carcinoma
		79238	GALE deficiency		
882	Fumarylacetoacetate hydrolase deficiency	79238	GALE-D	913	Gastrinoma

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2069	Gastrocutaneous syndrome	411777	GEKA	308487	Generalized GALE-D
2930	Gastrointestinal polyposis-ectodermal changes syndrome	26790	Gelatinous ascites	33355	Generalized hematopoietic hypoplasia
2930	Gastrointestinal polyposis-skin pigmentation-alopecia-fingernail changes syndrome	98957	Gelatinous drop-like corneal dystrophy	79402	Generalized junctional epidermolysis bullosa, non-Herlitz type
		2623	Geleophysic dwarfism		
		2623	Geleophysic dysplasia		
44890	Gastrointestinal stromal sarcoma	2073	Gélineau disease	329971	Generalized juvenile polyposis/juvenile polyposis coli
44890	Gastrointestinal stromal tumor	85448	Gelsolin amyloidosis		
2368	Gastroschisis	2074	Gemignani syndrome	167635	Generalized lichenoid papular eruption
355	Gaucher disease	251604	Gemistocytic astrocytoma		
77259	Gaucher disease type 1	2084	GEMSS syndrome	435628	Generalized lipodystrophy-progeroid features-severe intellectual disability syndrome
77260	Gaucher disease type 2	51608	Generalized arterial calcification of infancy		
77261	Gaucher disease type 3				
2072	Gaucher disease type 3C	79402	Generalized atrophic benign epidermolysis bullosa	141209	Generalized lymphatic anomaly
77261	Gaucher disease, subacute neuronopathic type	168632	Generalized basaloid follicular hamartoma syndrome	89842	Generalized mitis RDEB
2072	Gaucher disease-ophthalmoplegia-cardiovascular calcification syndrome			98806	Generalized cervical and upper-limb-onset dystonia
2072	Gaucher-like disease	528	Generalized congenital lipodystrophy	263543	Generalized peeling skin syndrome
308712	GBE deficiency, adult neuromuscular form			228429	Generalized congenital lipodystrophy type 4
308684	GBE deficiency, childhood combined hepatic and myopathic form	228429	Generalized congenital lipodystrophy with myopathy	263553	Generalized peeling skin syndrome type B
308698	GBE deficiency, childhood neuromuscular form			263558	Generalized deciduous skin
308670	GBE deficiency, congenital neuromuscular form	263548	Generalized deciduous skin type A	171876	Generalized pseudohypoadosteronism type 1
308655	GBE deficiency, fatal perinatal neuromuscular form	263553	Generalized deciduous skin type B	263543	Generalized PSS
		263558	Generalized deciduous skin type C	247353	Generalized pustular psoriasis
308638	GBE deficiency, non progressive hepatic form	231568	Generalized dominant dystrophic epidermolysis bullosa	3221	Generalized resistance to thyroid hormone
308621	GBE deficiency, progressive hepatic form	79399	Generalized EBS, non-Dowling-Meara type	308487	Generalized UDP-galactose-4-epimerase deficiency
360	GBM	99647	Generalized enchondromatosis with platyspondyly	308487	Generalized uridine diphosphate galactose-4-epimerase deficiency
98916	GBS, acute inflammatory demyelinating polyradiculoneuropathic form	79399	Generalized epidermolysis bullosa simplex, non-Dowling-Meara type	254704	Genetic hyperferritinemia without iron overload
329984	GCC	36387	Generalized epilepsy with febrile seizures-plus	99845	Genetic recurrent myoglobinuria
98962	GCD1	79137	Generalized epilepsy-paroxysmal dyskinesia syndrome	226316	Genetic transient congenital hypothyroidism
98963	GCD2			2075	Genito-palato-cardiac syndrome
25	GCDHD	308487	Generalized epimerase deficiency galactosemia	85201	Genitopatellar syndrome
98962	GCDI			2163	Genoa syndrome
98963	GCDII	157991	Generalized eruptive histiocytoma	85197	Genochondromatosis type 1
438274	GCGR-related hyperglucagonemia	157991	Generalized eruptive histiocytosis	93398	Genochondromatosis type 2
528	GCL	411777	Generalized eruptive keratoacanthoma	329813	Genome-wide paternal uniparental disomy mosaicism
228429	GCL4			1454	Gentile syndrome
2095	GCM syndrome	411777	Generalized eruptive keratoacanthomas of Grzybowski	217008	Genuine diffuse phlebectasia
380	GCPS	280774	Generalized essential telangiectasia	98961	Geographic corneal dystrophy
363976	GCT of bone			35686	Generalized exfoliative disease
98957	GDCD	36236	Generalized exfoliative disease	79137	GEPD
53697	GDD	1041	Generalized fetal edema		
366	GDE deficiency	308487	Generalized galactose epimerase deficiency	99095	Gerbode defect
324636	GDS			2808	Generalized GALE deficiency
36387	GEFS+	308487	Generalized GALE deficiency		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
213837	Germ cell cancer of cervix uteri	238763	Glaucoma secondary to spherophakia/ectopia lentis and megalocornea	51208	Glutamate formiminotransferase deficiency
213751	Germ cell cancer of corpus uteri			2195	Glutamate-aspartate transport defect
2077	German syndrome	2084	Glaucoma-ectopia-microspherophakia-stiff joints-short stature syndrome	33574	Glutamate-cysteine ligase deficiency
91352	Germinoma of the central nervous system			25	Glutaric acidemia type 1
2078	Geroderma osteodysplastica	2085	Glaucoma-sleep apnea syndrome	26791	Glutaric acidemia type 2
1117	Gershoni-Baruch-Leibo syndrome	354	GLB1 deficiency	35706	Glutaric acidemia type 3
221117	Gerstmann syndrome	360	Glioblastoma	25	Glutaric aciduria type 1
356	Gerstmann-Straussler-Scheinker syndrome	360	Glioblastoma multiforme	26791	Glutaric aciduria type 2
99926	Gestational choriocarcinoma	269197	Glioependymal/ependymal cyst	394532	Glutaric aciduria type 2, mild type
63275	Gestational pemphigoid	251582	Gliomatosis cerebri	394529	Glutaric aciduria type 2, severe neonatal type
280774	GET	251576	Gliosarcoma	35706	Glutaric aciduria type 3
84090	GFND	404476	Global developmental delay-lung cysts-overgrowth-Wilms tumor syndrome	25	Glutaryl-CoA dehydrogenase deficiency
314769	GH and PRL cosecreting pituitary adenoma			35706	Glutaryl-CoA oxidase deficiency
633	GH receptor deficiency	73223	Global developmental delay-osteopenia-ectodermal defect syndrome	25	Glutaryl-coenzyme A dehydrogenase deficiency
1802	Ghosal hematodiaphyseal dysplasia			32	Glutathione synthetase deficiency
1802	Ghosal syndrome	480898	Global developmental delay-visual anomalies-progressive cerebellar atrophy-truncal hypotonia syndrome	289846	Glutathione synthetase deficiency with 5-oxoprolinuria
83450	Ghost teeth			289849	Glutathione synthetase deficiency without 5-oxoprolinuria
314811	Ghrelin receptor deficiency	2791	Globodontia	33573	Glutathionuria
180267	Giant adenofibroma of the breast	487	Globoid cell leukodystrophy	284414	Glycerol kinase deficiency, adult form
643	Giant axonal neuropathy	83454	Glomangiomas	284408	Glycerol kinase deficiency, infantile form
397	Giant cell arteritis	→69735	Glomerulonephritis-sparse hair-telangiectasis syndrome	284411	Glycerol kinase deficiency, juvenile form
1190	Giant cell chondrodysplasia	84090	Glomerulopathy with fibronectin deposits	261476	Glycerol kinase deficiency-contiguous gene syndrome
251579	Giant cell glioblastoma	391651	Glomus tumor	255182	Glycine cleavage system L protein deficiency
139436	Giant cell histiocytomatosis	83454	Glomuvenous malformation	407	Glycine encephalopathy
363976	Giant cell tumor of bone	2616	Gloomy face syndrome	289891	Glycine N-methyltransferase deficiency
626	Giant congenital melanocytic nevus	141163	Glossopalatine ankylosis	365	Glycogen storage disease due to acid maltase deficiency
2494	Giant hypertrophic gastritis	221098	Glossopharyngeal neuralgia	308552	Glycogen storage disease due to acid maltase deficiency, infantile onset
626	Giant pigmented hairy nevus	221098	Glossosvopharyngeal neuralgia	420429	Glycogen storage disease due to acid maltase deficiency, late-onset
274	Giant platelet syndrome	404476	GLOW syndrome	57	Glycogen storage disease due to aldolase A deficiency
1065	Gillespie syndrome	255132	GLRX5-related sideroblastic anemia	364	Glycogen storage disease due to G6P deficiency
2025	Gingival fibromatosis-facial dysmorphism syndrome	97280	Glucagonoma	79258	Glycogen storage disease due to G6P deficiency type Ia
3473	Gingival fibromatosis-hepatosplenomegaly-other anomalies syndrome	97280	Glucagonoma syndrome	79259	Glycogen storage disease due to G6P deficiency type Ib
2026	Gingival fibromatosis-hypertrichosis syndrome	355	Glucocorticoid resistance	364	Glycogen storage disease due to glucose-6-phosphatase deficiency
2027	Gingival fibromatosis-progressive deafness syndrome	786	Glucocorticoid-remediable aldosteronism		
2709	Gingival hypertrophy-corneal dystrophy	403	Glucocorticoid-sensitive hypertension		
44890	GIST	403	Glucosamine N-acetyl-6-sulfatase deficiency		
97286	GIST-paranglioma dyad	79272	Glucose transporter type 1 deficiency		
358	Gitelman syndrome	71277	Glucose-galactose malabsorption		
3268	Giuffr�-Tsukahara syndrome	35710	Glucosidase 1 deficiency		
141209	GLA	79330	Glucosyltransferase 1 deficiency		
849	Glanzmann thrombasthenia	79320	Glucosyltransferase 2 deficiency		
666	Glass bone disease	79325	Glut1-DS		
1535	Glass-Chapman-Hockley syndrome	71277	Glut-1 deficiency Syndrome		
213833	Glassy cell carcinoma of the cervix uteri	71277	Glut-1 deficiency Syndrome		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
79258	Glycogen storage disease due to glucose-6-phosphatase deficiency type Ia	369	Glycogen storage disease due to liver glycogen phosphorylase deficiency	308621	Glycogen storage disease type 4, progressive hepatic form
				368	Glycogen storage disease type 5
79259	Glycogen storage disease due to glucose-6-phosphatase deficiency type Ib	2089	Glycogen storage disease due to liver glycogen synthase deficiency	369	Glycogen storage disease type 6
				371	Glycogen storage disease type 7
2088	Glycogen storage disease due to GLUT2 deficiency	264580	Glycogen storage disease due to liver phosphorylase kinase deficiency	264580	Glycogen storage disease type 9A
				79240	Glycogen storage disease type 9B
				264580	Glycogen storage disease type 9C
367	Glycogen storage disease due to glycogen branching enzyme deficiency	137625	Glycogen storage disease due to muscle and heart glycogen synthase deficiency	715	Glycogen storage disease type 9D
				715	Glycogen storage disease type 9E
308712	Glycogen storage disease due to glycogen branching enzyme deficiency, adult neuromuscular form	99849	Glycogen storage disease due to muscle beta-enolase deficiency	2088	Glycogen storage disease type 11
				284426	Glycogen storage disease type 11
				57	Glycogen storage disease type 12
308684	Glycogen storage disease due to glycogen branching enzyme deficiency, childhood combined hepatic and myopathic form	368	Glycogen storage disease due to muscle glycogen phosphorylase deficiency	→319646	Glycogen storage disease type 14
				263297	Glycogen storage disease type 15
				364	Glycogen storage disease type I
308698	Glycogen storage disease due to glycogen branching enzyme deficiency, childhood neuromuscular form	371	Glycogen storage disease due to muscle phosphofructokinase deficiency	79259	Glycogen storage disease type Ib
				365	Glycogen storage disease type II
				308552	Glycogen storage disease type II, infantile onset
308670	Glycogen storage disease due to glycogen branching enzyme deficiency, congenital neuromuscular form	715	Glycogen storage disease due to muscle phosphorylase kinase deficiency	420429	Glycogen storage disease type II, late onset
				→319646	Glycogen storage disease due to phosphoglucomutase deficiency
				366	Glycogen storage disease type III
308655	Glycogen storage disease due to glycogen branching enzyme deficiency, fatal perinatal neuromuscular form	713	Glycogen storage disease due to phosphoglycerate kinase 1 deficiency	367	Glycogen storage disease type IV
				308712	Glycogen storage disease type IV, adult neuromuscular form
				308684	Glycogen storage disease type IV, childhood combined hepatic and myopathic form
308638	Glycogen storage disease due to glycogen branching enzyme deficiency, non progressive hepatic form	97234	Glycogen storage disease due to phosphoglycerate mutase deficiency	308684	Glycogen storage disease type IV, childhood combined hepatic and myopathic form
				2089	Glycogen storage disease type 0a
				137625	Glycogen storage disease type 0b
308621	Glycogen storage disease due to glycogen branching enzyme deficiency, progressive hepatic form	364	Glycogen storage disease type 1	308698	Glycogen storage disease type IV, childhood neuromuscular form
				79258	Glycogen storage disease type 1a
				79259	Glycogen storage disease type 1b
308670	Glycogen storage disease due to glycogen branching enzyme deficiency, non progressive hepatic form	→79259	Glycogen storage disease type 1C	308670	Glycogen storage disease type IV, congenital neuromuscular form
				→79259	Glycogen storage disease type 1D
				365	Glycogen storage disease type 2
308655	Glycogen storage disease due to glycogen branching enzyme deficiency, progressive hepatic form	365	Glycogen storage disease type 2	308655	Glycogen storage disease type IV, fatal perinatal neuromuscular form
				308638	Glycogen storage disease type IV, non progressive hepatic form
				308621	Glycogen storage disease type IV, progressive hepatic form
366	Glycogen storage disease due to glycogen debranching enzyme deficiency	308552	Glycogen storage disease type 2, infantile onset	264580	Glycogen storage disease type IXa
				420429	Glycogen storage disease type 2, late onset
				79240	Glycogen storage disease type IXb
2089	Glycogen storage disease due to hepatic glycogen synthase deficiency	366	Glycogen storage disease type 3	264580	Glycogen storage disease type IXc
				367	Glycogen storage disease type 4
				264580	Glycogen storage disease type IXd
2364	Glycogen storage disease due to lactate dehydrogenase deficiency	308712	Glycogen storage disease type 4, adult neuromuscular form	715	Glycogen storage disease type IXe
				715	Glycogen storage disease type IXe
				368	Glycogen storage disease type V
284435	Glycogen storage disease due to lactate dehydrogenase H-subunit deficiency	308684	Glycogen storage disease type 4, childhood combined hepatic and myopathic form	369	Glycogen storage disease type VI
				371	Glycogen storage disease type VII
				2088	Glycogen storage disease type XI
284426	Glycogen storage disease due to lactate dehydrogenase M-subunit deficiency	308698	Glycogen storage disease type 4, childhood neuromuscular form	57	Glycogen storage disease type XII
				→319646	Glycogen storage disease type XIV
				263297	Glycogen storage disease type XV
34587	Glycogen storage disease due to LAMP-2 deficiency	308670	Glycogen storage disease type 4, congenital neuromuscular form	263297	Glycogen storage disease type XV
				263297	Glycogen storage disease with severe cardiomyopathy due to glycogenin deficiency
				308655	Glycogen storage disease type 4, fatal perinatal neuromuscular form
79240	Glycogen storage disease due to liver and muscle phosphorylase kinase deficiency	308638	Glycogen storage disease type 4, non progressive hepatic form		

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365	Glycogenosis due to acid maltase deficiency	264580	Glycogenosis due to liver phosphorylase kinase deficiency	263297	Glycogenosis type 15
308552	Glycogenosis due to acid maltase deficiency, infantile onset	137625	Glycogenosis due to muscle and heart glycogen synthase deficiency	364	Glycogenosis type I
57	Glycogenosis due to aldolase A deficiency	99849	Glycogenosis due to muscle beta-enolase deficiency	79258	Glycogenosis type Ia
79258	Glycogenosis due to glucose-6-phosphatase deficiency type 1a	368	Glycogenosis due to muscle glycogen phosphorylase deficiency	79259	Glycogenosis type Ib
79259	Glycogenosis due to glucose-6-phosphatase deficiency type 1b	371	Glycogenosis due to muscle phosphofruktokinase deficiency	365	Glycogenosis type II
79258	Glycogenosis due to glucose-6-phosphatase deficiency type Ia	715	Glycogenosis due to muscle phosphorylase kinase deficiency	308552	Glycogenosis type II, infantile onset
79259	Glycogenosis due to glucose-6-phosphatase transport defect type Ib	→319646	Glycogenosis due to phosphoglucomutase deficiency	420429	Glycogenosis type II, late onset
2088	Glycogenosis due to GLUT2 deficiency	713	Glycogenosis due to phosphoglycerate kinase 1 deficiency	366	Glycogenosis type III
367	Glycogenosis due to glycogen branching enzyme deficiency	97234	Glycogenosis due to phosphoglycerate mutase deficiency	367	Glycogenosis type IV
308712	Glycogenosis due to glycogen branching enzyme deficiency, adult neuromuscular form	2089	Glycogenosis type 0a	308712	Glycogenosis type IV, adult neuromuscular form
308684	Glycogenosis due to glycogen branching enzyme deficiency, childhood combined hepatic and myopathic form	137625	Glycogenosis type 0b	308684	Glycogenosis type IV, childhood combined hepatic and myopathic form
		364	Glycogenosis type 1	308698	Glycogenosis type IV, childhood neuromuscular form
		79259	Glycogenosis type 1b	308670	Glycogenosis type IV, congenital neuromuscular form
		365	Glycogenosis type 2	308655	Glycogenosis type IV, fatal perinatal neuromuscular form
308698	Glycogenosis due to glycogen branching enzyme deficiency, childhood neuromuscular form	308552	Glycogenosis type 2, infantile onset	308638	Glycogenosis type IV, non progressive hepatic form
		420429	Glycogenosis type 2, late onset	308621	Glycogenosis type IV, progressive hepatic form
		366	Glycogenosis type 3	264580	Glycogenosis type IXa
308670	Glycogenosis due to glycogen branching enzyme deficiency, congenital neuromuscular form	367	Glycogenosis type 4	79240	Glycogenosis type IXb
		308712	Glycogenosis type 4, adult neuromuscular form	264580	Glycogenosis type IXc
308655	Glycogenosis due to glycogen branching enzyme deficiency, fatal perinatal neuromuscular form	308684	Glycogenosis type 4, childhood combined hepatic and myopathic form	715	Glycogenosis type IXd
		308698	Glycogenosis type 4, childhood neuromuscular form	715	Glycogenosis type IXe
308638	Glycogenosis due to glycogen branching enzyme deficiency, non progressive hepatic form	308670	Glycogenosis type 4, congenital neuromuscular form	368	Glycogenosis type V
		308655	Glycogenosis type 4, fatal perinatal neuromuscular form	369	Glycogenosis type VI
308621	Glycogenosis due to glycogen branching enzyme deficiency, progressive hepatic form	308638	Glycogenosis type 4, non progressive hepatic form	371	Glycogenosis type VII
		308621	Glycogenosis type 4, progressive hepatic form	57	Glycogenosis type XII
366	Glycogenosis due to glycogen debranching enzyme deficiency	308621	Glycogenosis type 4, progressive hepatic form	→319646	Glycogenosis type XIV
2364	Glycogenosis due to lactate dehydrogenase deficiency	368	Glycogenosis type 5	263297	Glycogenosis type XV
284435	Glycogenosis due to lactate dehydrogenase H-subunit deficiency	369	Glycogenosis type 6	263297	Glycogenosis with severe cardiomyopathy due to glycogenin deficiency
		371	Glycogenosis type 7	93598	Glycolic aciduria
284426	Glycogenosis due to lactate dehydrogenase M-subunit deficiency	264580	Glycogenosis type 9A	354	GM1 gangliosidosis
		79240	Glycogenosis type 9B	79255	GM1 gangliosidosis type 1
34587	Glycogenosis due to LAMP-2 deficiency	264580	Glycogenosis type 9C	79256	GM1 gangliosidosis type 2
		715	Glycogenosis type 9D	79257	GM1 gangliosidosis type 3
79240	Glycogenosis due to liver and muscle phosphorylase kinase deficiency	715	Glycogenosis type 9E	796	GM2 gangliosidosis 0 variant
		284426	Glycogenosis type 11	309246	GM2 gangliosidosis, AB variant
		57	Glycogenosis type 12	309192	GM2 gangliosidosis, B variant, adult form
369	Glycogenosis due to liver glycogen phosphorylase deficiency	99849	Glycogenosis type 13	309178	GM2 gangliosidosis, B variant, infantile form
		→319646	Glycogenosis type 14	309185	GM2 gangliosidosis, B variant, juvenile form
				845	GM2 gangliosidosis, B, B1 variant
				309239	GM2 gangliosidosis, B1 variant
				101006	GM2 synthase deficiency
				626	GMN

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2090	GMS syndrome	377	Gorlin-Goltz syndrome	79477	GrisCELLi disease type 2
53697	Gnathodiaphyseal dysplasia	66629	GOSHS	79478	GrisCELLi disease type 3
602	GNE myopathy	280620	GOSR2-related progressive myoclonus ataxia	381	GrisCELLi-Pruniéras syndrome
100075	GNET			79476	GrisCELLi-Pruniéras syndrome type 1
481469	GNET type 1	2500	Gottron syndrome	79477	GrisCELLi-Pruniéras syndrome type 2
481475	GNET type 2	59135	Gowers disease	79478	GrisCELLi-Pruniéras syndrome type 3
481478	GNET type 3	900	GPA	2099	Grix-Blankenship-Peterson syndrome
481481	GNET type 4	280586	gPAPP deficiency		
79272	GNS deficiency	247353	GPP	3217	Groll-Hirschowitz syndrome
329984	Goblet cell adenocarcinoid	721	GPS	758	Gronblad-Strandberg-Touraine syndrome
329984	Goblet cell carcinoid	313808	GPSC		
329984	Goblet cell carcinoma	403	GRA	314613	Growing teratoma syndrome
329984	Goblet cell tumor	2763	Gracile bone dysplasia	391348	Growth and developmental delay-hypotonia-vision impairment-lactic acidosis syndrome
705	Goiter-deafness syndrome	53693	GRACILE syndrome		
373	Golabi-Rosen syndrome	39812	Graft versus host disease	→264200	Growth deficiency-brachydactyly-dysmorphism syndrome
351	Goldberg syndrome	505	Graham Little syndrome		
66629	Goldberg-Shprintzen megacolon syndrome	505	Graham Little-Piccardi-Lassueur syndrome	73273	Growth delay due to insulin-like growth factor I resistance
166272	Goldblatt chondrodysplasia	2111	Graham-Boyle-Troxell syndrome	73272	Growth delay due to insulin-like growth factor type 1 deficiency
166272	Goldblatt syndrome	52055	Graham-Cox syndrome		
3026	Goldblatt-Viljoen syndrome	→247691	Grand-Kaine-Fulling syndrome	2067	Growth delay-alopecia-pseudoanodontia-optic atrophy syndrome
2261	Goldblatt-Wallis syndrome	79094	Grange occlusive arterial syndrome		
374	Goldenhar syndrome	79094	Grange syndrome	3035	Growth delay-hydrocephaly-lung hypoplasia syndrome
53540	Goldmann-Favre syndrome	2097	Grant syndrome		
3032	Goldston syndrome	98962	Granular corneal dystrophy type 1	53693	Growth delay-aminoaciduria-cholestasis-iron overload-lactic acidosis-early death syndrome
1791	Gollop syndrome	98963	Granular corneal dystrophy type 2		
1986	Gollop-Wolfgang complex	98961	Granular corneal dystrophy type 3	73272	Growth delay-deafness- intellectual disability syndrome
2092	Goltz syndrome	98962	Granular corneal dystrophy type I		
2092	Goltz-Gorlin syndrome	98963	Granular corneal dystrophy type II	3035	Growth delay-hydrocephaly-lung hypoplasia syndrome
1532	Gómez-López-Hernández syndrome	98961	Granular corneal dystrophy type III		
206484	Gonadoblastoma	98963	Granular-lattice corneal dystrophy	314769	Growth hormone and prolactin cosecreting pituitary adenoma
432	Gonadotropic deficiency	86850	Granulocytic sarcoma		
759	Gonadotropin-dependant precocious puberty	900	Granulomatosis with polyangiitis	633	Growth hormone receptor deficiency
		183	Granulomatous allergic angiitis		
562	Gonadotropin-independent female-limited sexual precocity	64722	Granulomatous mastitis	97261	Growth hormone releasing factor tumor
		33111	Granulomatous slack skin		
2090	Goniodysgenesis-intellectual disability-short stature syndrome	99915	Granulosa cell cancer	53693	Growth restriction-aminoaciduria-cholestasis-iron overload-lactic acidosis-early death syndrome
		99915	Granulosa cell malignant tumor		
1482	Gonococcal conjunctivitis	35858	Gräsbeck-Imerslund disease	391366	Growth retardation-mild developmental delay-chronic hepatitis syndrome
3034	Gonzales-del Angel syndrome	69665	Gravidic intrahepatic cholestasis		
169105	Good syndrome	721	Gray platelet syndrome	2101	Grubben-de Cock-Borghgraef syndrome
1321	Goodman camptodactyly	293375	Grayson-Wilbrandt corneal dystrophy		
65798	Goodman syndrome			276405	Green jaundice
375	Goodpasture syndrome	99826	Green monkey disease		
75389	Goossens-Devriendt syndrome			1426	Greenberg dysplasia
757	Gordon hyperkalemia-hypertension syndrome	380	Greig cephalopolysyndactyly syndrome		
376	Gordon syndrome			495	Greither disease
1173	Gordon-Holmes syndrome	97261	GRF tumor		
73	Gorham disease			97261	GRFoma
73	Gorham syndrome	139474	Grisart-Destrée syndrome		
73	Gorham-Stout disease			381	GrisCELLi disease
377	Gorlin syndrome	79476	GrisCELLi disease type 1		
2095	Gorlin-Chaudhry-Moss syndrome				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
79259	GSD due to G6PT deficiency	713	GSD due to phosphoglycerate kinase 1 deficiency	368	GSD type V
2088	GSD due to GLUT2 deficiency			369	GSD type VI
367	GSD due to glycogen branching enzyme deficiency	97234	GSD due to phosphoglycerate mutase deficiency	371	GSD type VII
308712	GSD due to glycogen branching enzyme deficiency, adult neuromuscular form	2089	GSD type 0a	2088	GSD type XI
		137625	GSD type 0b	57	GSD type XII
308684	GSD due to glycogen branching enzyme deficiency, childhood combined hepatic and myopathic form	364	GSD type 1	263297	GSD type XV
		79259	GSD type 1 non a	263297	GSD with severe cardiomyopathy due to glycogenin deficiency
		79258	GSD type 1a	79258	GSDIa
		79259	GSD type 1b	79259	GSDIb
308698	GSD due to glycogen branching enzyme deficiency, childhood neuromuscular form	365	GSD type 2	366	GSDIII
		308552	GSD type 2, infantile onset	308712	GSDIV, adult neuromuscular form
308670	GSD due to glycogen branching enzyme deficiency, congenital neuromuscular form	420429	GSD type 2, late onset	308684	GSDIV, childhood combined hepatic and myopathic form
		366	GSD type 3	308698	GSDIV, childhood neuromuscular form
308655	GSD due to glycogen branching enzyme deficiency, fatal perinatal neuromuscular form	367	GSD type 4	308670	GSDIV, congenital neuromuscular form
		308712	GSD type 4, adult neuromuscular form	308655	GSDIV, fatal perinatal neuromuscular form
308638	GSD due to glycogen branching enzyme deficiency, non progressive hepatic form	308698	GSD type 4, childhood neuromuscular form	308638	GSDIV, non progressive hepatic form
308621	GSD due to glycogen branching enzyme deficiency, progressive hepatic form	308670	GSD type 4, congenital neuromuscular form	308621	GSDIV, progressive hepatic form
		308655	GSD type 4, fatal perinatal neuromuscular form	99849	GSDXIII
366	GSD due to glycogen debranching enzyme deficiency	308638	GSD type 4, non progressive hepatic form	→319646	GSDXIV
2089	GSD due to hepatic glycogen synthase deficiency	308621	GSD type 4, progressive hepatic form	2102	GTP cyclohydrolase I deficiency
2364	GSD due to lactate dehydrogenase deficiency	368	GSD type 5	98808	GTPCH1-deficient dopa-responsive dystonia
		369	GSD type 6	98808	GTPCH1-deficient DRD
284435	GSD due to lactate dehydrogenase H-subunit deficiency	371	GSD type 7	2102	GTPCH deficiency
284426	GSD due to lactate dehydrogenase M-subunit deficiency	264580	GSD type 9A	90020	Guam disease
		79240	GSD type 9B	319234	Guanarito hemorrhagic fever
34587	GSD due to LAMP-2 deficiency	264580	GSD type 9C	382	Guanidinoacetate methyltransferase deficiency
79240	GSD due to liver and muscle phosphorylase kinase deficiency	715	GSD type 9D	2785	Guibaud-Vainsel syndrome
		715	GSD type 9E	98916	Guillain-Barré syndrome, acute inflammatory demyelinating polyradiculoneuropathic form
369	GSD due to liver glycogen phosphorylase deficiency	97234	GSD type 10	231	Guinea worm disease
		2088	GSD type 11	1661	Guízar Vázquez-Luengas-Muñoz syndrome
264580	GSD due to liver phosphorylase kinase deficiency	284426	GSD type 11	2104	Guízar Vázquez-Sánchez-Manzano syndrome
		57	GSD type 12	1562	Gunal-Seber-Basaran syndrome
137625	GSD due to muscle and heart glycogen synthase deficiency	→319646	GSD type 14	79277	Günther disease
		263297	GSD type 15	1858	Gurrieri-Sammito-Bellussi syndrome
99849	GSD due to muscle beta-enolase deficiency	364	GSD type I	324561	Guttate hypopigmentation and punctate palmoplantar keratoderma
		79259	GSD type Ib	2957	Guttmacher syndrome
368	GSD due to muscle glycogen phosphorylase deficiency	365	GSD type II	39812	GVH
		308552	GSD type II, infantile onset	293375	GWCD
371	GSD due to muscle phosphofructokinase deficiency	420429	GSD type II, late onset	99914	Gynandroblastoma
		367	GSD type IV	414	Gyrate atrophy of choroid and retina
715	GSD due to muscle phosphorylase kinase deficiency	264580	GSD type IXa	168569	H syndrome
		79240	GSD type IXb		
→319646	GSD due to phosphoglucomutase deficiency	264580	GSD type IXc		
		715	GSD type IXd		
		715	GSD type IXe		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
139441	H-ABC	340	Hantavirus fever	157941	HDL1
2396	Haberland syndrome	319247	Hantavirus pulmonary syndrome	98934	HDL2
99803	Haddad syndrome	3294	Hapnes-Boman-Skeie syndrome	157946	HDL3
217026	Hadziselimovic syndrome	2967	Haptocorrin deficiency	98759	HDL4
91378	HAE	1490	Harboyan syndrome	313808	HDLS
100051	HAE 2	899	HARD syndrome	2237	HDR syndrome
100054	HAE 3	2812	Hard-skin syndrome, Parana type	402823	HDV
100050	HAE-I	85182	Hardcastle syndrome	288	HE
100051	HAE-II	1415	Hardikar syndrome	67037	Head and neck squamous cell carcinoma
100054	HAE-III	1177	Harding ataxia		
966	HAFF	457	Harlequin ichthyosis	254898	Hearing loss-encephaloneuropathy-obesity-valvulopathy syndrome
79263	Hagberg-Santavuori disease	199282	Harlequin syndrome		
2841	Hailey-Hailey disease	→216866	HARP syndrome	3225	Hearing loss-familial salivary gland insensitivity to aldosterone syndrome
2342	Haim-Munk syndrome	2115	Harrod syndrome		
1408	Hair defect-photosensitivity-intellectual disability syndrome	2116	Hartnup disease	1338	Heart defect-tongue hamartoma-polysyndactyly syndrome
		2116	Hartnup disorder		
69084	Hair-nail ectodermal dysplasia	2117	Hartsfield-Bixler-Demyer syndrome	1354	Heart defects-limb shortening syndrome
58017	Hairy cell leukemia	84085	HAS		
300878	Hairy cell leukemia variant	83601	Hashimoto encephalitis	875	Heart tumor of child
2220	Hairy elbows	99872	Hashimoto-Pritzker syndrome		
3387	Hairy throat syndrome	2994	Haspeslagh-Fryns-Muelenaere syndrome	392	Heart-hand syndrome type 1
955	Hajdu-Cheney syndrome			1350	Heart-hand syndrome type 2
2157	HAL deficiency	3325	HAT	1342	Heart-hand syndrome type 3
2985	Hal-Berg-Rudolph syndrome	2118	Hawkinsinuria	168796	Heart-hand syndrome, Slovenian type
2521	Halal syndrome	1071	Hay-Wells syndrome	1342	Heart-hand syndrome, Spanish type
1809	Halal-Setton-Wang syndrome	163596	Hb Bart's hydrops fetalis	1342	Heart-limb syndrome type 3
185	Halasz syndrome	231242	HbC-beta-thalassemia syndrome	442582	Heavy chain amyloidosis
138	Hall-Hittner syndrome	231249	HbE-beta-thalassemia syndrome	93556	Heavy chain deposition disease
2107	Hall-Riggs syndrome	93616	HbH disease	86864	Heavy chain disease
2108	Hallermann-Streiff syndrome	352657	HBID	2119	HEC syndrome
2109	Hallermann-Streiff-François syndrome, severe form	330032	HbLepore-beta-thalassemia syndrome	3377	Hecht syndrome
2109	Hallermann-Streiff-like syndrome	251359	HbS-beta-thalassemia syndrome	3377	Hecht-Beals syndrome
157850	Hallervorden-Spatz syndrome	251365	HbSC disease	2492	Hecht-Scott syndrome
2110	Hallux varus-preaxial polysyndactyly syndrome	251370	HbSD disease	238468	HED
		251375	HbSE disease	98813	HED-ID
3453	HAM syndrome	363412	HBSL	1882	HEDH syndrome
289326	HAM/TSP	88673	HCC	2787	Heide syndrome
314555	Hamamy syndrome	86864	HCD	3220	Heimler syndrome
2926	Hamanishi-Ueba-Tsuji syndrome	93556	HCDD	99932	Heiner syndrome
1217	Hamano-Tsukamoto syndrome	85458	HCHWA	178330	Heinz body anemia
2869	Hamartomatous intestinal polyposis	324723	HCHWA, Arctic type	86813	Helicoid peripapillary chorioretinal degeneration
93946	Hamel cerebro-palato-cardiac syndrome	100006	HCHWA, Dutch type	168782	Heller syndrome
		324718	HCHWA, Flemish type	244242	HELLP syndrome
79126	Hamman-Rich syndrome	100008	HCHWA, Icelandic type	1426	HEM dysplasia
73229	HANAC syndrome	324708	HCHWA, Iowa type	252054	Hemangioblastoma
1927	Hand and foot deformity-flat facies syndrome	324713	HCHWA, Italian type		
		324703	HCHWA, Piedmont type	2330	Hemangioma-thrombocytopenia syndrome
2438	Hand-foot-genital syndrome	100006	HCHWA-D		
2438	Hand-foot-uterus syndrome	58017	HCL		
99873	Hand-Schüller-Christian disease	300878	HCL-v	2128	Hemi 3 syndrome
989	Hanhart syndrome	163690	HCS	86908	Hemiconvulsion-hemiplegia-epilepsy syndrome
186	Hanot syndrome	306741	HD-HA syndrome		
340	Hantaviriosis	26106	HDGC		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2128	Hemicorporal hypertrophy			3325	Heparin-induced thrombocytopenia
443070	Hemicrania continua	248305	Hemolytic anemia due to glyceraldehyde-3-phosphate dehydrogenase deficiency	3325	Heparin-induced thrombocytopenia type 2
306741	Hemidystonia-hemiatrophy syndrome	35120	Hemolytic anemia due to pyrimidine 5' nucleotidase deficiency	156	Hepatic carnitine palmitoyl transferase 1 deficiency
1214	Hemifacial atrophy			156	Hepatic carnitine palmitoyl transferase I deficiency
1241	Hemifacial hyperplasia-strabismus syndrome	766	Hemolytic anemia due to red cell pyruvate kinase deficiency	386	Hepatic cystic hamartoma
141145	Hemifacial hypertrophy	275944	Hemolytic disease of the newborn with Kell alloimmunization	2031	Hepatic fibrosis-renal cysts-intellectual disability syndrome
141136	Hemifacial microsomia	90038	Hemolytic-uremic syndrome with diarrhea	369	Hepatic glycogen phosphorylase deficiency
2549	Hemifacial microsomia-radial defects syndrome	2134	Hemolytic-uremic syndrome without diarrhea	369	Hepatic phosphorylase deficiency
141148	Hemifacial myohyperplasia	93581	Hemolytic-uremic syndrome without diarrhea with anti-factor H antibodies	100035	Hepatic solitary necrotic tumor
276280	Hemihyperplasia-multiple lipomatosis syndrome	93578	Hemolytic-uremic syndrome without diarrhea with B factor anomaly	890	Hepatic veno-occlusive disease
99802	Hemimegalencephaly	93575	Hemolytic-uremic syndrome without diarrhea with C3 anomaly	79124	Hepatic veno-occlusive disease-immunodeficiency syndrome
306669	Hemiparkinsonism-hemiatrophy syndrome	357008	Hemolytic-uremic syndrome without diarrhea with DGKE deficiency	90073	Hepatitis B reinfection following liver transplantation
99050	Hemitruncus arteriosus	93579	Hemolytic-uremic syndrome without diarrhea with H factor anomaly	402823	Hepatitis D virus
139491	Hemochromatosis due to defect in ferroportin	93580	Hemolytic-uremic syndrome without diarrhea with I factor anomaly	402823	Hepatitis delta
79230	Hemochromatosis type 2			449	Hepatoblastoma
225123	Hemochromatosis type 3			54272	Hepatocellular adenoma
139491	Hemochromatosis type 4			88673	Hepatocellular carcinoma
447792	Hemochromatosis type 5			137681	Hepatoencephalopathy due to combined oxidative phosphorylation defect type 1
163596	Hemoglobin Bart's hydrops fetalis			137681	Hepatoencephalopathy due to COXPD1
2132	Hemoglobin C disease			95159	Hepatoerythropoietic porphyria
231242	Hemoglobin C-beta-thalassemia syndrome			905	Hepatolenticular degeneration
90039	Hemoglobin D disease			64743	Hepatoportal sclerosis
2133	Hemoglobin E disease			364	Hepatorenal glycogenosis
231249	Hemoglobin E-beta-thalassemia syndrome			882	Hepatorenal tyrosinemia
93616	Hemoglobin H disease			86882	Hepatosplenic T-cell lymphoma
330032	Hemoglobin Lepore-beta-thalassemia syndrome			306539	Hereditary acrokeratotic poikiloderma of Kindler-Weary
330041	Hemoglobin M disease			2907	Hereditary acrokeratotic poikiloderma, Weary type
280615	Hemoglobinopathy Toms River			447964	Hereditary adult-onset painful axonal polyneuropathy
244242	Hemolysis, elevated liver enzymes, low platelets in pregnancy			85450	Hereditary amyloid nephropathy
244242	Hemolysis-elevated liver enzymes-low platelets syndrome			93560	Hereditary amyloid nephropathy due to apolipoprotein A-I variant
86817	Hemolytic anemia due to adenylate kinase deficiency			238269	Hereditary amyloid nephropathy due to apolipoprotein A-II variant
714	Hemolytic anemia due to diphosphoglycerate mutase deficiency			93562	Hereditary amyloid nephropathy due to fibrinogen A alpha-chain variant
99138	Hemolytic anemia due to erythrocyte adenosine deaminase overproduction			93561	Hereditary amyloid nephropathy due to lysozyme variant
712	Hemolytic anemia due to glucophosphate isomerase deficiency			85450	Hereditary amyloidosis with primary renal involvement
90030	Hemolytic anemia due to glutathione reductase deficiency			85448	Hereditary amyloidosis, Finnish type
				228277	Hereditary anetoderma

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
91378	Hereditary angioedema	324708	Hereditary cerebral hemorrhage with amyloidosis, Iowa type	98873	Hereditary erythroblastic multinuclearity with a positive acidified-serum test (hempas)
100050	Hereditary angioedema type 1			36899	Hereditary essential myoclonus
100051	Hereditary angioedema type 2	324713	Hereditary cerebral hemorrhage with amyloidosis, Italian type	85195	Hereditary expansile polyostotic osteolytic dysplasia
100054	Hereditary angioedema type 3	324703	Hereditary cerebral hemorrhage with amyloidosis, Piedmont type	157846	Hereditary ferritinopathy
91378	Hereditary angioneurotic edema	48818	Hereditary ceruloplasmin deficiency	90045	Hereditary folate malabsorption
100050	Hereditary angioneurotic edema type 1	36382	Hereditary cervical artery dissection	469	Hereditary fructose intolerance
100051	Hereditary angioneurotic edema type 2	53372	Hereditary chin myoclonus	469	Hereditary fructose-1-phosphate aldolase deficiency
100054	Hereditary angioneurotic edema type 3	53372	Hereditary chin-trembling	469	Hereditary fructosemia
73229	Hereditary angiopathy-nephropathy-aneurysms-muscle cramps syndrome	676	Hereditary chronic pancreatitis	53372	Hereditary geniospasm
		422526	Hereditary clear cell renal cell adenocarcinoma	2024	Hereditary gingival fibromatosis
3115	Hereditary areflexic dystasia, Roussy-Lévy type	422526	Hereditary clear cell renal cell carcinoma	2024	Hereditary gingival hyperplasia
289601	Hereditary arterial and articular multiple calcification syndrome	293144	Hereditary clubfoot due to 5q31 microdeletion	774	Hereditary hemorrhagic telangiectasia
1416	Hereditary articular chondrocalcinosis	238578	Hereditary clubfoot due to 17q23.1-q23.2 microduplication	2604	Hereditary hollow visceral myopathy
1429	Hereditary benign chorea	293150	Hereditary clubfoot due to PITX1 point mutation	199285	Hereditary hypercarotenemia and vitamin A deficiency
352657	Hereditary benign corneal intraepithelial dyskeratosis	98434	Hereditary combined deficiency of factors II, VII, IX and X	238475	Hereditary hypercholanemia
352657	Hereditary benign intraepithelial dyskeratosis	98434	Hereditary combined deficiency of vitamin K-dependent clotting factors	3197	Hereditary hyperekplexia
91378	Hereditary bradykinine-induced angioedema	238722	Hereditary congenital controlateral synkinesia	3197	Hereditary hyperekplexia
221061	Hereditary brain cavernous angioma	238722	Hereditary congenital mirror movements	163	Hereditary hyperferritinemia with congenital cataracts
221061	Hereditary brain cavernous hemangioma			163	Hereditary hyperferritinemia-cataract syndrome
145	Hereditary breast and ovarian cancer syndrome	972	Hereditary continuous muscle fiber activity	2801	Hereditary hyperphosphatasia
227535	Hereditary breast cancer	79273	Hereditary coproporphria	157215	Hereditary hypophosphatemic rickets with hypercalciuria
227535	Hereditary breast carcinoma	60015	Hereditary cranium bifidum	55654	Hereditary hypotrichosis simplex
871	Hereditary bundle branch defect	168577	Hereditary cryohydrocytosis type 2	90368	Hereditary hypotrichosis simplex of the scalp
36382	Hereditary CAD	398088	Hereditary cryohydrocytosis with normal stomatin	217407	Hereditary hypotrichosis with recurrent skin vesicles
1416	Hereditary calcium pyrophosphate deposition	168577	Hereditary cryohydrocytosis with reduced stomatin	602	Hereditary inclusion body myopathy type 2
1416	Hereditary CC	98967	Hereditary crystalline stromal dystrophy of Schnyder	79091	Hereditary inclusion body myopathy type 3
30925	Hereditary CDI			324381	Hereditary inclusion body myopathy type 4
30925	Hereditary central diabetes insipidus	100008	Hereditary cystatin C amyloid angiopathy	178464	Hereditary inclusion body myopathy with early respiratory failure
221061	Hereditary cerebral cavernoma	26106	Hereditary diffuse cancer of stomach	79091	Hereditary inclusion body myopathy-joint contractures-ophthalmoplegia syndrome
221061	Hereditary cerebral cavernous malformation	26106	Hereditary diffuse gastric adenocarcinoma	300373	Hereditary infantile gigantism
85458	Hereditary cerebral hemorrhage with amyloidosis	26106	Hereditary diffuse gastric cancer	397692	Hereditary isolated aplastic anemia
324723	Hereditary cerebral hemorrhage with amyloidosis, Arctic type	313808	Hereditary diffuse leukoencephalopathy with axonal spheroids and pigmented glia	332	Hereditary juvenile megaloblastic anemia due to intrinsic factor deficiency
100006	Hereditary cerebral hemorrhage with amyloidosis, Dutch type	313808	Hereditary diffuse leukoencephalopathy with spheroids	2334	Hereditary keratitis
324718	Hereditary cerebral hemorrhage with amyloidosis, Flemish type	288	Hereditary elliptocytosis	493	Hereditary keratoacanthoma
100008	Hereditary cerebral hemorrhage with amyloidosis, Icelandic type	→247691	Hereditary endotheliopathy-retinopathy-nephropathy-stroke syndrome		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
411602	Hereditary late-onset Parkinson disease	640	Hereditary neuropathy with liability to pressure palsies	238269	Hereditary renal amyloidosis due to apolipoprotein A-II variant
523	Hereditary leiomyomatosis	279943	Hereditary neutrophilia	93562	Hereditary renal amyloidosis due to fibrinogen A alpha-chain variant
523	Hereditary leiomyomatosis and renal cell cancer	91378	Hereditary non histamine-induced angioedema	93561	Hereditary renal amyloidosis due to lysozyme variant
523	Hereditary leiomyomatosis with renal carcinoma	168583	Hereditary North American Indian childhood cirrhosis	94088	Hereditary renal hypouricemia
79452	Hereditary lymphedema type I	56	Hereditary ochronosis	788	Hereditary resistance to anti-vitamin K
90186	Hereditary lymphedema type II	30	Hereditary orotic aciduria	357027	Hereditary retinoblastoma
228277	Hereditary macular atrophy	98868	Hereditary ovalocytosis	221043	Hereditary sclerosing poikiloderma with tendon and pulmonary involvement
621	Hereditary methemoglobinemia	79141	Hereditary painful callosities	221039	Hereditary sclerosing poikiloderma, Weary type
157794	Hereditary mixed polyposis syndrome	86923	Hereditary palmoplantar hyperkeratosis, Gamborg-Nielsen type	280598	Hereditary sensorimotor neuropathy with hyperelastic skin
64748	Hereditary motor and sensory neuropathy type 3	86923	Hereditary palmoplantar keratoderma, Gamborg-Nielsen type	36386	Hereditary sensory and autonomic neuropathy type 1
773	Hereditary motor and sensory neuropathy type 4	47044	Hereditary papillary renal cell carcinoma	139564	Hereditary sensory and autonomic neuropathy type 1 with cough and gastroesophageal reflux
64751	Hereditary motor and sensory neuropathy type 5	99878	Hereditary parathyroids hyperplasia	139564	Hereditary sensory and autonomic neuropathy type 1B
90120	Hereditary motor and sensory neuropathy type 6	476102	Hereditary pediatric Behçet-like disease	970	Hereditary sensory and autonomic neuropathy type 2
64748	Hereditary motor and sensory neuropathy type III	168615	Hereditary persistence of alpha-fetoprotein	1764	Hereditary sensory and autonomic neuropathy type 3
90119	Hereditary motor and sensory neuropathy with acrodystrophy	46532	Hereditary persistence of fetal hemoglobin-beta-thalassemia syndrome	642	Hereditary sensory and autonomic neuropathy type 4
90103	Hereditary motor and sensory neuropathy with deafness, intellectual disability and absent sensory large myelinated fibers	251380	Hereditary persistence of fetal hemoglobin-sickle cell disease syndrome	64752	Hereditary sensory and autonomic neuropathy type 5
99950	Hereditary motor and sensory neuropathy, Lom type	29072	Hereditary pheochromocytoma-paraganglioma	314381	Hereditary sensory and autonomic neuropathy type 6
90117	Hereditary motor and sensory neuropathy, Okinawa type	300373	Hereditary pituitary hyperplasia	391397	Hereditary sensory and autonomic neuropathy type 7
90117	Hereditary motor and sensory neuropathy, proximal type	330061	Hereditary polymorphous light eruption of American Indians	478664	Hereditary sensory and autonomic neuropathy type 8
99953	Hereditary motor and sensory neuropathy, Russe Type	178345	Hereditary prepubertal gynecomastia	36386	Hereditary sensory and autonomic neuropathy type I
1839	Hereditary mucoepithelial dysplasia	828	Hereditary progressive arthroophthalmopathy	139564	Hereditary sensory and autonomic neuropathy type IB
171723	Hereditary mucosal leukokeratosis	98808	Hereditary progressive dystonia with marked diurnal fluctuation	970	Hereditary sensory and autonomic neuropathy type II
136	Hereditary multi-infarct dementia	158025	Hereditary progressive mucinous histiocytosis	1764	Hereditary sensory and autonomic neuropathy type III
→3460	Hereditary multicentric osteolysis	178464	Hereditary proximal myopathy with early respiratory failure	642	Hereditary sensory and autonomic neuropathy type IV
523	Hereditary multiple cutaneous leiomyomas	264675	Hereditary pulmonary alveolar proteinosis	64752	Hereditary sensory and autonomic neuropathy type V
83454	Hereditary multiple glomangiomas	440427	Hereditary pulmonary alveolar proteinosis with hepatic involvement	314381	Hereditary sensory and autonomic neuropathy type VI
2590	Hereditary myoclonus-progressive distal muscular atrophy syndrome	275777	Hereditary pulmonary arterial hypertension	391397	Hereditary sensory and autonomic neuropathy type VII
43115	Hereditary myopathy with lactic acidosis due to ISCU deficiency	→288	Hereditary pyropoikilocytosis	478664	Hereditary sensory and autonomic neuropathy type VIII
1062	Hereditary neurocutaneous malformation	85450	Hereditary renal amyloidosis		
456333	Hereditary neuroendocrine tumor of small bowel	93560	Hereditary renal amyloidosis due to apolipoprotein A-I variant		
456333	Hereditary neuroendocrine tumor of small intestine				
30925	Hereditary neurogenic diabetes insipidus				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
139573	Hereditary sensory and autonomic neuropathy with deafness and global delay	93160	Hereditary vitamin D-resistant rickets	309246	Hexosaminidase activator deficiency
		903	Hereditary von Willebrand disease	796	Hexosaminidases A and B deficiency
391397	Hereditary sensory and autonomic neuropathy with hyperhidrosis and gastrointestinal dysfunction	98805	Hereditary whispering dysphonia	309169	Hexosaminidases A and B deficiency, adult form
		170	Hereditary woolly hair syndrome	309155	Hexosaminidases A and B deficiency, infantile form
139578	Hereditary sensory and autonomic neuropathy with spastic paraplegia	170	Hereditary woolly hair syndrome		
		3467	Hereditary xanthinuria		
456318	Hereditary sensory neuropathy-deafness-dementia syndrome	3202	Hereditary xerocytosis	1041	HF
		773	Hereditary ataxia polyneuritisformis	2438	HFGS
456318	Hereditary sensory neuropathy-sensorineural hearing loss-dementia syndrome	275777	Hereditary pulmonary arterial hypertension	2744	HGPPS
		3411	Herlyn-Werner syndrome	740	HGPS
213524	Hereditary site-specific ovarian cancer syndrome	79430	Hermansky-Pudlak syndrome	79271	HGSNAT deficiency
		183678	Hermansky-Pudlak syndrome type 2	163	HHCS
100996	Hereditary spastic paraparesis type 15	231531	Hermansky-Pudlak syndrome type 7	86908	HHE syndrome
		231537	Hermansky-Pudlak syndrome type 8	415	HHH syndrome
822	Hereditary spherocytosis	280663	Hermansky-Pudlak syndrome type 9	276280	HHML
84093	Hereditary thermosensitive neuropathy	183678	Hermansky-Pudlak syndrome with neutropenia	157215	HHRH
71493	Hereditary thrombocytopenia with early-onset myelofibrosis			774	HHT
480851	Hereditary thrombocytopenia with normal platelets	231500	Hermansky-Pudlak syndrome with pulmonary fibrosis	457	HI
				231512	Hermansky-Pudlak syndrome without pulmonary fibrosis
268322	Hereditary thrombocytopenia with normal platelets	2139	Hernández-Aguirre Negrete syndrome	35878	HI/HA syndrome
				2786	Hernández-Fragoso syndrome
71290	Hereditary thrombocytopenia with normal platelets-hematological cancer predisposition syndrome	→247691	HERNS syndrome	602	HIBM2
		1930	Herpes simplex encephalitis	79091	HIBM3
329319	Hereditary thrombocytopenia with transverse limb defect	1930	Herpes simplex meningo-encephalitis	324381	HIBM4
				1808	Hidrotic ectodermal dysplasia, Christianson-Fourie type
82	Hereditary thrombophilia due to congenital antithrombin 3 deficiency	1930	Herpes simplex neuroinvasion	189	Hidrotic ectodermal dysplasia
				1930	Herpetiform pemphigus
217467	Hereditary thrombophilia due to congenital histidine-rich (poly-L) glycoprotein deficiency	208524	Herpetiform pemphigus	343	HIDS
		369	Hers disease	137577	HIE
217467	Hereditary thrombophilia due to congenital HRG deficiency	1486	Herva disease	137577	HIE
		314970	HES-L	330012	High altitude pulmonary edema
745	Hereditary thrombophilia due to congenital protein C deficiency	314950	HES-M	171201	High anorectal malformation
		314950	HES-N	314029	High bone mass OI
743	Hereditary thrombophilia due to congenital protein S deficiency	314962	HES-R	314029	High bone mass osteogenesis imperfecta
		640	Heterozygous microdeletion 17p11.2p12	480541	High grade B-cell lymphoma with MYC and/ or BCL2 and/or BCL6 rearrangement
745	Hereditary thrombophilia due to PC deficiency	3450	Heterozygous OSMED	363396	High myopia-sensorineural deafness syndrome
		3450	Heterozygous otospondylomegapiphysal dysplasia	3181	High scapula
205	Hereditary unconjugated hyperbilirubinemia	845	Hexosaminidase A deficiency	231080	High-grade dysplasia in patients with Barrett esophagus
79234	Hereditary unconjugated hyperbilirubinemia type 1	309192	Hexosaminidase A deficiency, adult form	251646	High-grade ependymoma
79235	Hereditary unconjugated hyperbilirubinemia type 2	309239	Hexosaminidase A deficiency, B1 variant	213777	High-grade neuroendocrine carcinoma of the cervix uteri
→247691	Hereditary vascular retinopathy	309178	Hexosaminidase A deficiency, infantile form	213731	High-grade neuroendocrine carcinoma of the corpus uteri
→247691	Hereditary vascular retinopathy-Raynaud phenomenon-migraine syndrome	309185	Hexosaminidase A deficiency, juvenile form	213777	High-grade neuroendocrine carcinoma of the uterine cervix

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213731	High-grade neuroendocrine carcinoma of the uterine corpus	2158	Histidinuria-renal tubular defect syndrome	2117	Holoprosencephaly-ectrodactyly-cleft lip palate syndrome
101088	HIGM1	50918	Histiocytic necrotizing lymphadenitis	2570	Holoprosencephaly-fetal akinesia/hypokinesia sequence syndrome
101089	HIGM2				
101090	HIGM3				
101091	HIGM4				
101092	HIGM5				
183663	HIGM with susceptibility to opportunistic infections	3325	HIT	2166	Holoprosencephaly-postaxial polydactyly syndrome
		→138	Hittner-Hirsch-Kreh syndrome		
		443291	HIV-associated cancer		
183666	HIGM without susceptibility to opportunistic infections	443291	HIV-related cancer	3186	Holoprosencephaly-radial heart renal anomalies syndrome
		1573	HJMD	392	Holt-Oram syndrome
99978	Hilar CCA	572	HLA class 2-negative severe combined immunodeficiency	2167	Holzgreve-Wagner-Rehder syndrome
99978	Hilar cholangiocarcinoma				
84085	Hinman syndrome				
84085	Hinman-Allen syndrome	2248	HLHS	30924	HOMG1
1164	Hinson-Pepys disease	412	HLP type 3	34528	HOMG2
2114	Hip dysplasia, Beukes type	523	HLRCC	31043	HOMG3
3408	Hip dysplasia-enchondromata-ecchondroma syndrome	2213	HMC syndrome	2168	Homocarnosinase deficiency
		178464	HMERF	1361	Homocarnosinosis
411593	Hirata disease	20	HMG-CoA lyase deficiency	2168	Homocarnosinosis
65684	Hirayama disease	35701	HMG-CoA synthase deficiency	394	Homocystinuria due to cystathionine beta-synthase deficiency
388	Hirschsprung disease	157794	HMPS		
261537	Hirschsprung disease and intellectual disability due to 2q22 microdeletion	64748	HMSN 3	395	Homocystinuria due to methylene tetrahydrofolate reductase deficiency
		773	HMSN 4		
		64751	HMSN 5		
261552	Hirschsprung disease and intellectual disability due to a ZEB2 point mutation	401964	HMSN2 with giant axons	622	Homocystinuria without methylmalonic aciduria
		64748	HMSN III		
261537	Hirschsprung disease and intellectual disability due to del(2)(q22)	90119	HMSN with acrodystrophy	56	Homogentisic acid oxidase deficiency
		99950	HMSN, Lom type	163596	Homozygous alpha0-thalassemia
		99950	HMSN-Lom		
		90117	HMSNP		
261537	Hirschsprung disease and intellectual disability due to monosomy 2q22	99953	HMSNR	391665	Homozygous familial hypercholesterolemia
		69084	HNED	14	Homozygous familial hypobetalipoproteinemia
		93111	HNFB-MODY		
2155	Hirschsprung disease-deafness-polydactyly syndrome	640	HNPP	→288	Homozygous hereditary elliptocytosis
2151	Hirschsprung disease-ganglioneuroblastoma syndrome	67037	HNSCC	98958	Honey-droplet corneal dystrophy
		1979	Hoepffner-Dreyer-Reimers syndrome	98960	Honeycomb corneal dystrophy
2152	Hirschsprung disease-intellectual disability syndrome	2349	Hoffman syndrome	78	Hookworm infection
				391665	HoFH
2153	Hirschsprung disease-nail hypoplasia-dysmorphism syndrome	414	HOGA	2744	Horizontal gaze palsy with progressive scoliosis
		454718	Holmes-Adie syndrome	397	Horton disease
2150	Hirschsprung disease-type D brachydactyly syndrome	→994	Holmes-Benacerraf syndrome	392	HOS
		3328	Holmes-Collins syndrome	166412	Hot water reflex epilepsy
2026	Hirsutism-congenital gingival hyperplasia syndrome	93970	Holmes-Gang syndrome	1352	Houlston-Irton-Temple syndrome
2156	Hirsutism-skeletal dysplasia-intellectual disability syndrome	2143	Holmes-Schepens syndrome	99907	House allergic alveolitis
		79242	Holocarboxylase synthetase deficiency	2198	Howell-Evans syndrome
3283	His bundle tachycardia	2162	Holoprosencephaly	3322	Hoyeraal-Hreidarsson syndrome
2157	HIS deficiency	2165	Holoprosencephaly-caudal dysgenesis syndrome	306669	HP-HA syndrome
2157	Histidase deficiency			275777	HPAH
2157	Histidine ammonia-lyase deficiency			98808	Holoprosencephaly-craniosynostosis syndrome
2157	Histidinemia				
2157	Histidinuria	2163			HPE

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46532	HPFH-beta-thalassemia syndrome	1930	HSV encephalitis	→263463	Humerospinal dysostosis
251380	HPFH-sickle cell disease syndrome	285	HT-EDS	3383	Humerus trochlea aplasia
247262	HPMR	289326	HTLV-1-associated myelopathy/tropical spastic paraparesis	580	Hunter syndrome
436	HPP			217085	Hunter syndrome type A
293958	HPPD			217093	Hunter syndrome type B
47044	HPRCC	482077	HTRA1-related autosomal dominant cerebral angiopathy	→35069	Hunter-Carpenter-McDonald syndrome
79233	HPRT1 partial deficiency	482077	HTRA1-related autosomal dominant cerebral small vessel disease	2715	Hunter-Jurenka-Thompson syndrome
510	HPRT complete deficiency				
510	HPRT deficiency grade IV	228116	Hughes-Stovin syndrome	97340	Hunter-McAlpine craniosynostosis
79233	HPRT deficiency, grade I	438279	Human infection by orthopoxvirus	3365	Hunter-Rudd-Hoffmann syndrome
79233	HPRT partial deficiency				
79233	HPRT-related gout	289326	Human T-lymphotropic virus type I-associated myelopathy/tropical spastic paraparesis	1390	Hunter-Thompson-Reed syndrome
79233	HPRT-related hyperuricemia			399	Huntington chorea
79430	HPS			399	Huntington disease
183678	HPS2	289326	Human T-lymphotropic virus type-1-associated myelopathy/tropical spastic paraparesis	401901	Huntington disease phenocopy due to C9ORF72 expansions
231531	HPS7				
231537	HPS8	294973	Humeral agenesis/hypoplasia	157941	Huntington disease-like 1
280663	HPS9	295063	Humeral agenesis/hypoplasia, bilateral	98934	Huntington disease-like 2
231500	HPS with pulmonary fibrosis				
231512	HPS without pulmonary fibrosis	295061	Humeral agenesis/hypoplasia, unilateral	157946	Huntington disease-like 3
99880	HPT-JT				
2323	HRD syndrome	294973	Humeral intercalary meromelia	98759	Huntington disease-like 4
84085	HS	295063	Humeral intercalary meromelia, bilateral	401901	Huntington disease-like syndrome due to C9ORF72 expansions
36386	HSAN1				
139564	HSAN1B	295061	Humeral intercalary meromelia, unilateral	363694	HUPRA syndrome
456318	HSAN1E				
970	HSAN2	3265	Humero-radial fusion	384	Huriez syndrome
1764	HSAN3	295211	Humero-radial fusion, bilateral	93473	Hurler disease
642	HSAN4	295209	Humero-radial fusion, unilateral	93473	Hurler syndrome
64752	HSAN5	3265	Humero-radial synostosis	93476	Hurler-Scheie syndrome
314381	HSAN6	295211	Humero-radial synostosis, bilateral	330061	Hutchinson summer prurigo
391397	HSAN7	295209	Humero-radial synostosis, unilateral	740	Hutchinson-Gilford progeria syndrome
478664	HSAN8	3266	Humero-radio-ulnar fusion	93160	HVDRR
139564	HSAN with cough and gastroesophageal reflux	295207	Humero-radio-ulnar fusion, bilateral	364039	HVLL
		295205	Humero-radio-ulnar fusion, unilateral	→247691	HVR
139573	HSAN with deafness and global delay	294975	Humero-radio-ulnar intercalary transverse meromelia	53698	Hyaline body myopathy
391397	HSAN with hyperhidrosis and gastrointestinal dysfunction				
139578	HSAN with spastic paraplegia	295087	Humero-radio-ulnar intercalary transverse meromelia, bilateral	70587	Hyaline membrane disease
2182	HSAS	295085	Humero-radio-ulnar intercalary transverse meromelia, unilateral	530	Hyalinosis cutis et mucosae
388	HSCR				
391417	HSD10 deficiency	3266	Humero-radio-ulnar synostosis	67041	Hyaluronidase deficiency
85295	HSD10 deficiency, atypical type	295207	Humero-radio-ulnar synostosis, bilateral	400	Hydatid disease
391428	HSD10 deficiency, classic type				
391428	HSD10 deficiency, infantile type	295205	Humero-radio-ulnar synostosis, unilateral	99927	Hydatidiform mole
391457	HSD10 deficiency, neonatal type				
391417	HSD10 disease	→263463	Humero-spinal dysostosis	400	Hydatidosis
85295	HSD10 disease, atypical type	94056	Humero-ulnar fusion	2898	Hyde Forster-McCarthy-Berry syndrome
391428	HSD10 disease, classic type	295215	Humero-ulnar fusion, bilateral	2177	Hydranencephaly
391428	HSD10 disease, infantile type	295213	Humero-ulnar fusion, unilateral	330021	Hydrargyria
391457	HSD10 disease, neonatal type	94056	Humero-ulnar synostosis	330061	Hydroa aestivale
30924	HSH				
456318	HSN1E	295215	Humero-ulnar synostosis, bilateral	330058	Hydroa vacciniforme
		295213	Humero-ulnar synostosis, unilateral	364039	Hydroa vacciniforme-like lymphoma
				2182	Hydrocephalus with stenosis of the aqueduct of Sylvius
				899	Hydrocephalus-agyria-retinal dysplasia syndrome
				2186	Hydrocephalus-blue sclerae-nephropathy syndrome

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916	Hydrocephalus-cleft palate-joint contractures syndrome	168588	Hyperandrogenism due to cortisone reductase deficiency	276556	Hyperinsulinemic hypoglycemia due to UCP2 deficiency
2180	Hydrocephalus-costovertebral dysplasia-Sprengel anomaly syndrome	90	Hyperargininemia	79299	Hyperinsulinism due to glucokinase deficiency
		234	Hyperbilirubinemia type 2	71212	Hyperinsulinism due to glutamodehydrogenase deficiency
		3111	Hyperbilirubinemia, Rotor type		
2119	Hydrocephalus-endocardial fibroelastosis-cataract syndrome	276405	Hyperbiliverdinemia	324575	Hyperinsulinism due to HNF1A deficiency
		306661	Hypercalcemic tumoral calcinosis	263455	Hyperinsulinism due to HNF4A deficiency
2183	Hydrocephalus-obesity-hypogonadism syndrome	2196	Hypercalciuria-bilateral macular coloboma syndrome		
1397	Hydrocephaly-cerebellar agenesis syndrome	209902	Hypercholesterolemia due to cholesterol 7alpha-hydroxylase deficiency	263458	Hyperinsulinism due to INSR deficiency
2184	Hydrocephaly-low insertion umbilicus syndrome				
2181	Hydrocephaly-tall stature-joint laxity syndrome	83639	Hypercoagulability syndrome due to glycosylphosphatidylinositol deficiency	165991	Hyperinsulinism due to monocarboxylate transporter 1 deficiency
221126	Hydrocephaly/hydranencephaly due to cerebral vasculopathy	1032	Hyperdibasic aminoaciduria type 1	71212	Hyperinsulinism due to SCHAD deficiency
		470	Hyperdibasic aminoaciduria type 2		
2189	Hydroletharus	3197	Hyperkplexia	71212	Hyperinsulinism due to short chain 3-hydroxylacyl-CoA dehydrogenase deficiency
2473	Hydrometrocolpos-postaxial polydactyly syndrome	163985	Hyperkplexia-epilepsy syndrome		
		408	Hyperglycerolemia	165991	Hyperinsulinism due to SLC16A1 deficiency
2704	Hydronephrosis-inverted smile syndrome	2410	Hypergonadotropic hypogonadism-cataract syndrome	276556	Hyperinsulinism due to UCP2 deficiency
1041	Hydrops fetalis	243	Hypergonadotropic ovarian dysgenesis	35878	Hyperinsulinism-hyperammonemia syndrome
1426	Hydrops-ectopic calcification-motheaten syndrome	2157	Hyperhistidinemia		
20	Hydroxymethylglutaric aciduria	742	Hyperimidodipeptiduria	757	Hyperkalemia-hypertension syndrome, Gordon type
401	Hymenolepiasis	343	Hyperimmunoglobulinemia D with recurrent fever		
309147	Hyper-beta-alaninemia	2314	Hyperimmunoglobulin E syndrome type 1	682	Hyperkalemic periodic paralysis
343	Hyper-IgD syndrome			682	Hyperkalemic PP
101090	Hyper-IgM syndrome due to CD40 deficiency	2314	Hyperimmunoglobulin E-recurrent infection syndrome	409	Hyperkeratosis lenticularis perstans
101088	Hyper-IgM syndrome due to CD40 ligand deficiency			1662	Hyperkeratosis-contracture syndrome
101088	Hyper-IgM syndrome due to CD40L deficiency	343	Hyperimmunoglobulinemia D syndrome	1336	Hyperkeratosis-hyperpigmentation syndrome
101092	Hyper-IgM syndrome due to UNG deficiency	343	Hyperimmunoglobulinemia D with periodic fever	682	HyperKPP
101092	Hyper-IgM syndrome due to uracil N-glycosylase	79299	Hyperinsulinemic hypoglycemia due to glucokinase deficiency	140905	Hyperlipidemia due to hepatic triglyceride lipase deficiency
101088	Hyper-IgM syndrome type 1	324575	Hyperinsulinemic hypoglycemia due to HNF1A deficiency	412	Hyperlipidemia type 3
101089	Hyper-IgM syndrome type 2			→444490	Hyperlipoproteinemia type 1
101090	Hyper-IgM syndrome type 3	263455	Hyperinsulinemic hypoglycemia due to HNF4A deficiency	412	Hyperlipoproteinemia type 3
101091	Hyper-IgM syndrome type 4	263458	Hyperinsulinemic hypoglycemia due to INSR deficiency	70470	Hyperlipoproteinemia type 5
101092	Hyper-IgM syndrome type 5			2203	Hyperlysinemia
183663	Hyper-IgM syndrome with susceptibility to opportunistic infections	263458	Hyperinsulinemic hypoglycemia due to insulin receptor deficiency	2203	Hyperlysinemia type I
				3124	Hyperlysinemia type II
183666	Hyper-IgM syndrome without susceptibility to opportunistic infections	276603	Hyperinsulinemic hypoglycemia due to Kir6.2 deficiency, diazoxide-resistant focal form	289891	Hypermethioninemia due to glycine N-methyltransferase deficiency
				289891	Hypermethioninemia due to GNMT deficiency
309147	Hyperalaninemia	71212	Hyperinsulinemic hypoglycemia due to short chain 3-hydroxylacyl-CoA dehydrogenase deficiency	88618	Hypermethioninemia due to S-adenosylhomocysteine hydrolase deficiency
927	Hyperammonemia due to N-acetylglutamate synthase deficiency	276598	Hyperinsulinemic hypoglycemia due to SUR1 deficiency, diazoxide-resistant focal form	289290	Hypermethioninemia encephalopathy due to adenosine kinase deficiency
401948	Hyperammonemic encephalopathy due to carbonic anhydrase VA deficiency				

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289290	Hypermethioninemia encephalopathy due to ADK deficiency	93604	Hyperprostaglandin E syndrome	251523	Hyperzincemia and hypercalprotectinemia
		889	Hypersensitivity angitis		
		1519	Hypertelorism, Teebi type		
73267	Hypernychthemeral syndrome	2211	Hypertelorism-hypospadias-polysyndactyly syndrome	2435	Hypo- and hypermelanotic cutaneous macules-retarded growth-intellectual disability syndrome
414	Hyperornithinemia				
414	Hyperornithinemia-gyrate atrophy of choroid and retina syndrome				
415	Hyperornithinemia-hyperammonemia-homocitrullinuria syndrome	2213	Hypertelorism-microtia-facial clefting syndrome	289157	Hypocalcemic vitamin D-dependent rickets
2801	Hyperostosis corticalis deformans juvenilis	2745	Hypertelorism-oesophageal abnormality-hypospadias syndrome		
		293958	Hypertelorism-preauricular sinus-punctual pits-deafness syndrome		
3416	Hyperostosis corticalis generalisata	293958	Hypertelorism-preauricular sinus-punctual pits-hearing loss syndrome	93160	Hypocalcemic vitamin D-resistant rickets
443098	Hyperostosis cranialis interna	88660	Hypertension due to gain-of-function mutations in the mineralocorticoid receptor	100032	Hypocalcified amelogenesis imperfecta
77296	Hyperostosis frontalis interna			93297	Hypochondrogenesis
2780	Hyperostosis generalisata with striations			429	Hypochondroplasia
99880	Hyperparathyroidism-jaw tumor syndrome	757	Hypertensive hyperkalemia	36412	Vasculocomplementemic urticarial vasculitis
295002	Hyperphalangy	423	Hyperthermia of anesthesia	2228	Hypodontia-dysplasia of nails syndrome
295140	Hyperphalangy in digits 2-5	2220	Hypertrichosis cubiti-short stature syndrome	2228	Hypodontia-nail dysgenesis syndrome
295142	Hyperphalangy, bilateral	2222	Hypertrichosis lanuginosa congenita	185	Hypogenetic lung syndrome
295140	Hyperphalangy, unilateral	2222	Hypertrichosis universalis	989	Hypoglossia-hypodactyly syndrome
1388	Hyperphalangy-clinodactyly of index finger with Pierre Robin syndrome	2026	Hypertrichosis with or without gingival hyperplasia	→261483	Hypogonadism-gynecomastia-X-linked intellectual disability syndrome
238583	Hyperphenylalaninemia	966	Hypertrichosis-acromegaloid facial appearance syndrome		
13	Hyperphenylalaninemia due to 6-pyruvoyltetrahydropterin synthase deficiency	966	Hypertrichosis-acromegaloid facial features syndrome	2233	Hypogonadism-mitral valve prolapse-intellectual disability syndrome
238583	Hyperphenylalaninemia due to BH4 deficiency	1231	Hypertrichosis-atrophic skin-ectropion-macrostromia syndrome	141333	Hypogonadism-short stature-coloboma-preaxial polydactyly syndrome
1578	Hyperphenylalaninemia due to dehydratase deficiency	966	Hypertrichosis-coarse face syndrome		
226	Hyperphenylalaninemia due to dihydropteridine reductase deficiency	319182	Hypertrichosis-short stature-facial dysmorphism-developmental delay syndrome	2230	Hypogonadotropic hypogonadism-frontoparietal alopecia syndrome
2102	Hyperphenylalaninemia due to GTP cyclohydrolase deficiency	1517	Hypertrichotic osteochondrodysplasia, Cantu type	2235	Hypogonadotropic hypogonadism-retinitis pigmentosa syndrome
1578	Hyperphenylalaninemia due to pterin-4-alpha-carbinolamine dehydratase deficiency	324525	Hypertrophic cardiomyopathy and renal tubular disease due to mitochondrial DNA mutation	293967	Hypogonadotropic hypogonadism-severe microcephaly-sensorineural deafness-dysmorphism syndrome
238583	Hyperphenylalaninemia due to tetrahydrobiopterin deficiency	324525	Hypertrophic cardiomyopathy and renal tubular disease due to mtDNA mutation	293967	Hypogonadotropic hypogonadism-severe microcephaly-sensorineural hearing loss-dysmorphism syndrome
1578	Hyperphenylalaninemia with primapterinuria	217601	Hypertrophic cardiomyopathy due to intensive athletic training	363523	Hypohidrosis-enamel hypoplasia-palmoplantar keratoderma-intellectual disability syndrome
2209	Hyperphenylalaninemic embryopathy	329883	Hypertrophic gastropathy without hypoproteinemia	238468	Hypohidrotic ectodermal dysplasia
3416	Hyperphosphatasemia tarda	90282	Hypertrophic or verrucous lupus erythematosus	98813	Hypohidrotic ectodermal dysplasia with immunodeficiency
247262	Hyperphosphatasia-intellectual disability syndrome	2224	Hypertryptophanemia	1882	Hypohidrotic ectodermal dysplasia-hypothyroidism-ciliary dyskinesia syndrome
→79189	Hyperpipecolatemia	217330	Hyperuricemia-anemia-renal failure syndrome	293964	Hypoinsulinemic hypoglycemia and body hemihypertrophy
157798	Hyperplastic polyposis syndrome				
682	HyperPP	363694	Hyperuricemia-pulmonary hypertension-renal failure-alkalosis syndrome	681	Hypokalemic periodic paralysis
419	Hyperprolinemia type 1				
79101	Hyperprolinemia type 2				
				30924	Hypomagnesemia caused by selective magnesium malabsorption

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
30924	Hypomagnesemia intestinal type 1	→3157	Hypopituitarism-postaxial polydactyly syndrome	163690	Hypotonia-cystinuria syndrome
1790	Hypomandibular faciocranial dysostosis	99058	Hypoplasia of the mitral valve annulus	79507	Hypotonia-failure to thrive-microcephaly syndrome
100033	Hypomaturation amelogenesis imperfecta	722	Hypoplasminogenemia	371364	Hypotonia-speech impairment-severe cognitive delay syndrome
100034	Hypomaturation-hypoplastic amelogenesis imperfecta with taurodontism	100031	Hypoplastic amelogenesis imperfecta	55654	Hypotrichosis simplex
435	Hypomelanosis of Ito	2248	Hypoplastic left heart syndrome	90368	Hypotrichosis simplex of the scalp
2680	Hypomyelination neuropathy-arthrogryposis syndrome	293864	Hypoplastic pancreas-intestinal atresia-hypoplastic gallbladder syndrome	1573	Hypotrichosis with juvenile macular degeneration
139441	Hypomyelination with atrophy of basal ganglia and cerebellum	3332	Hypoplastic tibiae-postaxial polydactyly syndrome	1573	Hypotrichosis with juvenile macular dystrophy
363412	Hypomyelination with brain stem and spinal cord involvement and leg spasticity	→216866	Hypoprebetalipoproteinemia-acanthocytosis-retinitis pigmentosa-pallidal degeneration syndrome	444	Hypotrichosis, Marie Unna type
447893	Hypomyelination-cerebellar atrophy-hypoplasia of the corpus callosum syndrome	327	Hypoproconvertinemia	91132	Hypotrichosis-congenital ichthyosis syndrome
85163	Hypomyelination-congenital cataract syndrome	2494	Hypoproteinemic hypertrophic gastropathy	330029	Hypotrichosis-deafness syndrome
88637	Hypomyelination-hypogonadotropic hypogonadism-hypodontia syndrome	325	Hypoprothrombinemia	2266	Hypotrichosis-intellectual disability, Lopes type
3453	Hypoparathyroidism-Addison disease-mucocutaneous candidiasis syndrome	2250	Hyposmia-nasal and ocular hypoplasia-hypogonadotropic hypogonadism syndrome	69735	Hypotrichosis-lymphedema-telangiectasia-membranoproliferative glomerulonephritis syndrome
2237	Hypoparathyroidism-deafness-renal disease syndrome	2745	Hypospadias-dysphagia syndrome	69735	Hypotrichosis-lymphedema-telangiectasia-renal defect syndrome
2323	Hypoparathyroidism-intellectual disability-dysmorphism syndrome	2745	Hypospadias-hypertelorism syndrome	307936	Hypotrichosis-osteolysis-periodontitis-palmoplantar hyperkeratosis syndrome
2323	Hypoparathyroidism-short stature-intellectual disability-seizures syndrome	→1299	Hypospadias-hypertelorism-coloboma and deafness syndrome	307936	Hypotrichosis-osteolysis-periodontitis-palmoplantar keratoderma syndrome
436	Hypophosphatasia	2261	Hypospadias-intellectual disability, Goldblatt type syndrome	307936	Hypotrichosis-striate palmoplantar hyperkeratosis-acroosteolysis-periodontitis syndrome
314621	Hypophyseal duplication	2353	Hypotelorism-cleft palate-hypospadias syndrome	307936	Hypotrichosis-striate palmoplantar keratoderma-acroosteolysis-periodontitis syndrome
99725	Hypophyseal gigantism	443101	Hypothalamic adipic hypernatraemia syndrome	79233	Hypoxanthine guanine phosphoribosyltransferase 1 partial deficiency
324561	Hypopigmentation and punctate keratosis of the palms and soles	672	Hypothalamic hamartoblastoma syndrome	510	Hypoxanthine guanine phosphoribosyltransferase complete deficiency
42665	Hypopigmentation-deafness syndrome	86906	Hypothalamic hamartomas with gelastic seizures	79233	Hypoxanthine guanine phosphoribosyltransferase deficiency, grade I
79477	Hypopigmentation-immunodeficiency with or without neurologic impairment syndrome	→3157	Hypothalamic insufficiency-secondary microcephaly-visual impairment-urinary anomalies syndrome	510	Hypoxanthine guanine phosphoribosyltransferase deficiency, grade IV
79476	Hypopigmentation-neurologic impairment syndrome	226307	Hypothyroidism due to deficient transcription factors involved in pituitary development or function	79233	Hypoxanthine guanine phosphoribosyltransferase partial deficiency
324561	Hypopigmentation-punctate palmoplantar keratoderma syndrome	90673	Hypothyroidism due to TSH receptor mutations	137577	Hypoxic and ischemic brain injury in the newborn
91354	Hypopituitarism due to empty sella turcica syndrome	1226	Hypothyroidism-cleft palate syndrome	137577	Hypoxic-ischemic encephalopathy
→3157	Hypopituitarism-micropenis-cleft lip/palate syndrome	3047	Hypothyroidism-dysmorphism-postaxial polydactyly-intellectual disability syndrome	682	HYPP
→3157	Hypopituitarism-microphthalmia syndrome	91131	Hypotonia and ichthyosis due to dolichol phosphate deficiency	63440	Hypsicephaly
		137908	Hypotonia with lactic acidemia and hyperammonemia		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
63440	Hypocephaly	2272	Ichthyosis-oral and digital anomalies syndrome	228315	Idiopathic hypersomnia with long sleep time
576	I-cell disease				
480512	IAD	363992	Ichthyosis-short stature-brachydactyly-microspherophakia syndrome	228318	Idiopathic hypersomnia without long sleep time
724	IAEP			449427	Idiopathic hypertrophic pachymeningitis
158048	IAHS	289347	IDH	1572	Idiopathic immunoglobulin deficiency
293168	IAHSP	3306	idic(15)	51608	Idiopathic infantile arterial calcification
254509	Iatrogenic botulism	930	Idiopathic achalasia	238624	Idiopathic intracranial hypertension
95619	Iatrogenic or traumatic pituitary deficiency	930	Idiopathic achalasia of esophagus	85193	Idiopathic juvenile osteoporosis
363424	IBA57 deficiency	724	Idiopathic acute eosinophilic pneumonia	247234	Idiopathic late-onset cerebellar ataxia
→33364	IBIDS syndrome	139423	Idiopathic acute transverse myelitis	314017	Idiopathic linear interstitial keratitis
611	IBM	480512	Idiopathic adult ductopenia	33577	Idiopathic lobular panniculitis
602	IBM2	422	Idiopathic and/or familial pulmonary arterial hypertension	90158	Idiopathic localized lipodystrophy
79091	IBM3	280914	Idiopathic anterior uveitis	353344	Idiopathic macular telangiectasia type 1
52430	IBMPFD	88	Idiopathic aplastic anemia	353351	Idiopathic macular telangiectasia type 3
1576	IBSN	399307	Idiopathic avascular necrosis	84065	Idiopathic malabsorption due to bile acid synthesis defects
31709	ICCA syndrome	399307	Idiopathic AVN	73	Idiopathic massive osteolysis
64734	ICE syndrome	1980	Idiopathic basal ganglia calcification	97560	Idiopathic membranous glomerulonephritis
2268	ICF syndrome	171684	Idiopathic basal ganglia calcification	2573	Idiopathic Moyamoya disease
455	Ichthyosis bullosa of Siemens	84065	Idiopathic bilateral vestibulopathy	2774	Idiopathic multicentric osteolysis with or without nephropathy
457	Ichthyosis congenita, Harlequin type	88	Idiopathic bile acid malabsorption	824	Idiopathic myelofibrosis
289586	Ichthyosis exfoliativa	60033	Idiopathic bone marrow failure	45452	Idiopathic neonatal atrial flutter
457	Ichthyosis fetalis, Harlequin type	1320	Idiopathic bronchiectasis	33577	Idiopathic nodular panniculitis
2273	Ichthyosis follicularis-alopecia-photophobia syndrome	1320	Idiopathic camptocormia	51608	Idiopathic obliterative arteriopathy
2273	Ichthyosis follicularis-atrichia-photophobia syndrome	188	Idiopathic camptocormism	441	Idiopathic orthostatic hypotension
79504	Ichthyosis hystrix gravior	163703	Idiopathic capillary leak syndrome	280921	Idiopathic panuveitis
79503	Ichthyosis hystrix of Curth-Macklin	228000	Idiopathic catastrophic epileptic encephalopathy	747	Idiopathic PAP
79503	Ichthyosis hystrix, Curth-Macklin type	169615	Idiopathic CD4 lymphocytopenia	480524	Idiopathic peliosis hepatis
88621	Ichthyosis prematurity syndrome	2902	Idiopathic central precocious puberty	480524	Idiopathic peliosis hepatitis
281190	Ichthyosis variegata	95717	Idiopathic chronic eosinophilic pneumonia	444316	Idiopathic phalangeal acroosteolysis
281190	Ichthyosis with confetti	209919	Idiopathic congenital hypothyroidism	444316	Idiopathic phalangeal acroosteolysis
79504	Ichthyosis, Lambert type	35062	Idiopathic copper-associated cirrhosis	280917	Idiopathic posterior uveitis
2269	Ichthyosis-alopecia-eclabion-ectropion-intellectual disability syndrome	35062	Idiopathic disseminated CMV infection	1320	Idiopathic progressive lumbar kyphosis
2267	Ichthyosis-cheek-eyebrow syndrome	447881	Idiopathic disseminated cytomegalovirus infection	747	Idiopathic pulmonary alveolar proteinosis
91132	Ichthyosis-follicular atrophoderma-hypotrichosis syndrome	480512	Idiopathic dropped head syndrome	275766	Idiopathic pulmonary arterial hypertension
91132	Ichthyosis-follicular atrophoderma-hypotrichosis-hypohidrosis syndrome	256	Idiopathic ductopenia	1676	Idiopathic pulmonary artery dilatation
2274	Ichthyosis-hepatosplenomegaly-cerebellar degeneration syndrome	247724	Idiopathic dystonia	2032	Idiopathic pulmonary fibrosis
91132	Ichthyosis-hypotrichosis syndrome	329874	Idiopathic eosinophilic myositis	99931	Idiopathic pulmonary hemosiderosis
59303	Ichthyosis-hypotrichosis-sclerosing cholangitis syndrome	64722	Idiopathic giant cell myocarditis	35061	Idiopathic recurrent and disabling cutaneous herpes
2278	Ichthyosis-intellectual disability-dwarfism-renal impairment syndrome	86908	Idiopathic granulomatous mastitis	251307	Idiopathic recurrent pericarditis
→1643	Ichthyosis-male hypogonadism syndrome	2197	Idiopathic hemiconvulsion-hemiplegia syndrome		
		3260	Idiopathic hypercalciuria		
		33208	Idiopathic hypereosinophilic syndrome		
			Idiopathic hypersomnia		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
276174	Idiopathic recurrent stupor	449566	IgG4-related eosinophilic angiocentric fibrosis	37042	Immune dysregulation-polyendocrinopathy-enteropathy-X-linked syndrome
251307	Idiopathic relapsing pericarditis				
40923	Idiopathic retinal perivasculitis	90003	IgG4-related hepatopathy	364013	Immune fetal edema
40923	Idiopathic retinal vasculitis	449395	IgG4-related kidney disease		
209943	Idiopathic retinal vasculitis-aneurysms-neuroretinitis syndrome	63999	IgG4-related mediastinitis	364013	Immune fetal hydrops
		238593	IgG4-related mesenteritis	364013	Immune HF
49041	Idiopathic retroperitoneal fibrosis	449563	IgG4-related ophthalmic disease	364013	Immune hydrops fetalis
458718	Idiopathic SCAD	449427	IgG4-related pachymeningitis	206569	Immune myopathy with myocyte necrosis
35065	Idiopathic severe pneumococemia	280302	IgG4-related pancreatitis		
458718	Idiopathic spontaneous coronary artery dissection	449400	IgG4-related periaortitis	1959	Immune pancytopenia
		49041	IgG4-related retroperitoneal fibrosis	3002	Immune thrombocytopenia
69061	Idiopathic steroid-sensitive nephrotic syndrome	447764	IgG4-related sclerosing cholangitis	3002	Immune thrombocytopenic purpura
		449432	IgG4-related sialadenitis	206569	Immune-mediated necrotizing myopathy
93209	Idiopathic steroid-sensitive nephrotic syndrome with diffuse mesangial proliferation	449432	IgG4-related submandibular gland disease		
		64744	IgG4-related thyroid disease	86886	Immunoblastic lymphadenopathy
93206	Idiopathic steroid-sensitive nephrotic syndrome with focal segmental glomerulosclerosis	183675	IgG subclass deficiency with IgA subclass deficiency	34592	Immunodeficiency by defective expression of HLA class 1
		329235	IGSF1 deficiency syndrome	572	Immunodeficiency by defective expression of HLA class 2
93206	Idiopathic steroid-sensitive nephrotic syndrome with focal segmental hyalinosis	364013	IHF		
		86908	IHHS		
93207	Idiopathic steroid-sensitive nephrotic syndrome with minimal change	371364	IHPRF syndrome	169150	Immunodeficiency due to a C5 to C9 component complement deficiency
		91132	IHS		
		59303	IHSC		
99858	Idiopathic syringomyelia	238624	IIH	169147	Immunodeficiency due to a classical component pathway complement deficiency
256	Idiopathic torsion dystonia	85193	IJO		
98806	Idiopathic torsion dystonia of mixed type	238569	IL10-related early-onset IBD	169147	Immunodeficiency due to an early component of complement deficiency
		238569	IL10-related early-onset inflammatory bowel disease		
3347	Idiopathic tracheobronchomegaly	477661	IL21-related infantile IBD	169150	Immunodeficiency due to a late component of complement deficiency
209956	Idiopathic uveal effusion syndrome	477661	IL21-related infantile inflammatory bowel disease		
130	Idiopathic ventricular fibrillation, Brugada type	100078	Ileal neuroendocrine neoplasm	169147	Immunodeficiency due to CD25 deficiency
228140	Idiopathic ventricular fibrillation, non Brugada type	100078	Ileal neuroendocrine tumor		
280384	IDMDC	238621	Ileal pouch anal anastomosis related faecal incontinence	169100	Immunodeficiency due to ficolin3 deficiency
580	Iduronate 2-sulfatase deficiency	1150	Illum syndrome	331190	Immunodeficiency due to interleukin-1 receptor-associated kinase-4 deficiency
217085	Iduronate 2-sulfatase deficiency type A	79466	ILVEN		
217093	Iduronate 2-sulfatase deficiency type B	85173	IMAGe syndrome	70592	Immunodeficiency due to factor H anomaly
		247718	IMAM		
92050	IED	42062	Iminoglycinuria	331187	Immunodeficiency due to selective anti-polysaccharide antibody deficiency
91132	IFAH syndrome	284362	Immature interstitial mesenchymal tumor		
2273	IFAP syndrome	398987	Immature teratoma of ovary	70593	Immunodeficiency due to MASP-2 deficiency
332	IFD	289465	Immigration delay disease		
329903	Ig-mediated membranoproliferative glomerulonephritis	→244	Immotile cilia syndrome, Kartagener type	200421	Immunodeficiency due to selective anti-polysaccharide antibody deficiency
329903	Ig-mediated MPGN	2901	Immune brachial plexus neuropathy	200418	Immunodeficiency with factor H anomaly
761	IgA vasculitis	169090	Immune dysfunction due to T-cell inactivation due to calcium entry defect		
329874	IGCM	238569	Immune dysregulation-inflammatory bowel disease-arthritis-recurrent infections syndrome	2268	Immunodeficiency with factor I anomaly
79099	IGDA			2268	Immunodeficiency-centromeric instability-facial anomalies syndrome
73272	IGF-1 deficiency	238569	Immune dysregulation-inflammatory bowel disease-arthritis-recurrent infections syndrome	647	Immunodeficiency-microcephaly-chromosomal instability syndrome
449400	IgG4-related aortitis				
79078	IgG4-related dacryoadenitis and sialadenitis				

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935	Immunodeficiency-short limb dwarfism syndrome	178478	Infant intestinal toxemia botulism	1928	Infantile lobar hyperinflation
761	Immunoglobulin A vasculitis	178478	Infant intestinal toxin-mediated botulism	667	Infantile malignant osteopetrosis
169110	Immunoglobulin heavy chain deficiency	70587	Infant respiratory distress syndrome	247165	Infantile mercury intoxication
329903	Immunoglobulin-mediated membranoproliferative glomerulonephritis	178487	Infant-like botulism	247165	Infantile mercury poisoning
329903	Immunoglobulin-mediated MPGN	247165	Infantile acrodynia	456312	Infantile multisystem neurologic-endocrine-pancreatic disease
100025	Immunoproliferative small intestinal disease	99749	Infantile agranulocytosis	2591	Infantile myofibromatosis
97567	Immunotactoid glomerulonephritis	99725	Infantile and juvenile forms of acromegaly	79263	Infantile NCL
97567	Immunotactoid glomerulopathy	70590	Infantile apnea	93591	Infantile nephronophthisis
456312	IMNEPD	51608	Infantile arteriosclerosis	35069	Infantile neuroaxonal dystrophy
206569	IMNM	2679	Infantile axonal neuropathy	79263	Infantile neuronal ceroid lipofuscinosis
857	Imperforate anus with hand, foot and ear anomalies	89938	Infantile Bartter syndrome with sensorineural deafness	289860	Infantile NKH
2759	Imperforate oropharynx-costo vertebral anomalies syndrome	1576	Infantile bilateral striatal necrosis	289860	Infantile non-ketotic hyperglycinemia
71276	Imploding antrum syndrome	178478	Infantile botulism	251304	Infantile onset panniculitis with uveitis and systemic granulomatosis
35069	INAD	314911	Infantile Canavan disease	1186	Infantile onset spinocerebellar ataxia
35069	INAD1	137675	Infantile cardiomyopathy with histiocytoid change	67047	Infantile optic atrophy with chorea and spastic paraplegia
254509	Inadvertent botulism	217557	Infantile cellular interstitial pneumonitis	85179	Infantile osteopetrosis with neuroaxonal dysplasia
45453	Incessant infant ventricular tachycardia	313850	Infantile cerebellar-retinal degeneration	247651	Infantile phosphoethanolaminuria
79263	INCL	402364	Infantile cerebral and cerebellar atrophy with postnatal progressive microcephaly	247651	Infantile Rathburn disease
231226	Inclusion body beta-thalassemia	77260	Infantile cerebral Gaucher disease	772	Infantile Refsum disease
199267	Inclusion body fibromatosis	1313	Infantile choroidocerebral calcification syndrome	254864	Infantile reversible cytochrome C oxidase deficiency myopathy
602	Inclusion body myopathy type 2	31709	Infantile convulsions and choreoathetosis	263410	Infantile spasms-psychomotor retardation-progressive brain atrophy-basal ganglia disease syndrome
79091	Inclusion body myopathy type 3	1310	Infantile cortical hyperostosis	3451	Infantile spasms
52430	Inclusion body myopathy with Paget disease of bone and frontotemporal dementia	199267	Infantile digital fibromatosis	3173	Infantile spasms-broad thumbs syndrome
611	Inclusion body myositis	87876	Infantile dysmorphic sialidosis	83330	Infantile spinal muscular atrophy
254693	Incomplete hydatidiform mole	238455	Infantile dystonia-parkinsonism	1576	Infantile striatonigral degeneration
254693	Incomplete molar pregnancy	364063	Infantile epileptic-dyskinetic encephalopathy	1576	Infantile striatonigral necrosis
157769	Incomplete situs inversus	300373	Infantile gigantism due to pituitary hyperplasia	1575	Infantile striatothalamic degeneration
180079	Incomplete unilateral aplasia of the Müllerian ducts	289860	Infantile glycine encephalopathy	255241	Infantile subacute necrotizing encephalopathy with leukodystrophy
180079	Incomplete unilateral Müllerian aplasia	79255	Infantile GM1 gangliosidosis	255249	Infantile subacute necrotizing encephalopathy with nephrotic syndrome
464	Incontinentia pigmenti	309155	Infantile GM2 gangliosidosis 0 variant	3311	Infantile symmetrical thalamic degeneration
435	Incontinentia pigmenti type 1	293603	Infantile hereditary endothelial dystrophy	2176	Infantile systemic hyalinosis
158019	Indeterminate cell histiocytosis	352563	Infantile hypertrophic cardiomyopathy due to MRPL44 deficiency	1577	Infantile thalamic degeneration
1388	Index finger anomaly-Pierre Robin syndrome	247651	Infantile hypophosphatasia	2768	Infantile tibia vara
98848	Indolent systemic mastocytosis	371364	Infantile hypotonia-psychomotor retardation-characteristic facies syndrome	137675	Infantile xanthomatous cardiomyopathy
1909	Indomethacin embryofetopathy	79076	Infantile juvenile polyposis syndrome	293168	Infantile-onset ascending hereditary spastic paralysis
70587	Infant acute respiratory distress syndrome	206436	Infantile Krabbe disease		
70587	Infant ARDS				
178478	Infant botulism				
1943	Infant epilepsy with migrant focal crisis				
178478	Infant intestinal botulism				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
284332	Infantile-onset autosomal recessive nonprogressive cerebellar ataxia	319465	Inherited AML	364577	Intellectual disability-brachydactyly-Pierre Robin syndrome
457205	Infantile-onset axonal motor and sensory neuropathy-optic atrophy-neurodegenerative syndrome	319462	Inherited cancer-predisposing syndrome due to biallelic BRCA2 mutations	→324737	Intellectual disability-cataract-coloboma-kyphosis syndrome
		282166	Inherited CJD	3042	Intellectual disability-cataracts-calcified pinnae-myopathy syndrome
391316	Infantile-onset mesial temporal lobe epilepsy with severe cognitive regression	210141	Inherited congenital spastic quadriplegia		171860
1451	Infantile-onset multisystem inflammatory disease	210141	Inherited congenital spastic tetraplegia	397709	Intellectual disability-coarse face-macrocephaly-cerebellar hypoplasia syndrome
		282166	Inherited Creutzfeldt-Jakob disease		Intellectual disability-coarse face-macrocephaly-cerebellar hypotrophy syndrome
171714	Infantile-onset symptomatic epilepsy syndrome-developmental stagnation-blindness syndrome	859	Inherited deficiency of transcobalamin	397709	Intellectual disability-coarse face-macrocephaly-cerebellar hypotrophy syndrome
1145	Infantile-onset X-linked spinal muscular atrophy	100054	Inherited estrogen-associated angioedema		
		100054	Inherited estrogen-associated angioneurotic edema		
781	Infection due to Coxiella burnetii	100054	Inherited estrogen-dependent angioedema	3454	Intellectual disability-developmental delay-contractures syndrome
279922	Infectious anterior uveitis	100054	Inherited estrogen-dependent angioedema		
137593	Infectious epithelial keratitis			100054	Inherited estrogen-dependent angioneurotic edema
279925	Infectious panuveitis	71278	Inherited glutamine synthetase deficiency		
279919	Infectious posterior uveitis				
289347	Infective dermatitis associated with HTLV-1	71278	Inherited GS deficiency	→280	Intellectual disability-dysmorphism-intrauterine growth retardation syndrome
289347	Infective dermatitis associated with human T-lymphotropic virus type 1	289548	Inherited isolated adrenal insufficiency due to partial CYP11A1 deficiency		
289347	Infective dermatitis associated with human T-lymphotropic virus type I	225968	Inherited predisposition to essential thrombocythemia	171851	Intellectual disability-enteropathy-deafness-peripheral neuropathy-ichthyosis-keratoderma syndrome
99123	Inferior caval vein interruption				
155889	Inferior palpebral coloboma	37	Inherited zinc deficiency	2139	Intellectual disability-epilepsy-bulbous nose syndrome
99123	Inferior vena cava interruption				
280794	Infiltrative small vesicular DCM	63259	Iniensephaly	127	Intellectual disability-epilepsy-endocrine disorders syndrome
280794	Infiltrative small vesicular diffuse cutaneous mastocytosis	178475	Inoculation botulism		
		411593	Insulin autoimmune syndrome	468620	Intellectual disability-epilepsy-extrapyramidal syndrome
85445	Inflammatory amyloidosis	2297	Insulin-resistance syndrome type A		
79466	Inflammatory linear verrucous epidermal nevus	2298	Insulin-resistance syndrome type B		
		97279	Insulinoma	404440	Intellectual disability-facial dysmorphism syndrome due to SETD5 haploinsufficiency
178342	Inflammatory myofibroblastic tumor	100973	Intellectual disability associated with fragile site FRAXE		
160148	Inflammatory myoglandular polyps	464311	Intellectual disability syndrome due to a DYRK1A point mutation		
247718	Inflammatory myopathy with abundant macrophages	166108	Intellectual disability, Birk-Barel type	404440	Intellectual disability-facial dysmorphism syndrome due to SETD5 haploinsufficiency
263553	Inflammatory peeling skin syndrome	3079	Intellectual disability, Buenos-Aires type		
48918	Inflammatory pseudotumor of skeletal muscle	→324737	Intellectual disability, Kahrizi type	363611	Intellectual disability-feeding difficulties-developmental delay-microcephaly syndrome
90003	Inflammatory pseudotumor of the liver	2557	Intellectual disability, Mietens-Weber type		
238305	Infundibulo-neurohypophysitis	3080	Intellectual disability, Wolff type	369847	Intellectual disability-hyperkinetic movement-truncal ataxia syndrome
95513	Infundibulo-panhypophysitis	289483	Intellectual disability-alacrima-achalasia syndrome		
1849	Infundibulopelvic stenosis-multicystic kidney syndrome			2466	Intellectual disability-aphasia-shuffling gait-adducted thumbs syndrome
247257	Inhalation anthrax disease	3041	Intellectual disability-balding-patella luxation-acromicria syndrome		
254504	Inhalation botulism				
247257	Inhalational anthrax				
254504	Inhalational botulism				
319465	Inherited acute myeloid leukemia				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
314575	Intellectual disability-hypotonia-brachycephaly-pyloric stenosis-cryptorchidism syndrome	363528	Intellectual disability-strabismus syndrome	440392	Interstitial lung disease due to SP-C deficiency
166108	Intellectual disability-hypotonia-facial dysmorphism syndrome	397941	Intellectual disability-truncal obesity syndrome	440392	Interstitial lung disease due to surfactant protein C deficiency
3050	Intellectual disability-hypotonia-skin hyperpigmentation syndrome	75858	Intellectual disability-truncal obesity-retinal dystrophy-micropenis syndrome	99092	Interventricular septum aneurysm
356996	Intellectual disability-hypotonia-spasticity-sleep disorder syndrome	1478	Interatrial communication	1201	Intestinal atresia type IIIb
3451	Intellectual disability-hypsarrhythmia syndrome	1478	Interauricular communication	178481	Intestinal botulism
436151	Intellectual disability-loss of expressive language-facial dysmorphism syndrome	51890	Intercostal nerve syndrome	178481	Intestinal colonization botulism
457279	Intellectual disability-macrocephaly-hypotonia-behavioral abnormalities syndrome	86900	Interdigitating cell sarcoma	92050	Intestinal epithelial dysplasia
3067	Intellectual disability-microcephaly-phalangeal-facial abnormalities syndrome	86900	Interdigitating dendritic cell sarcoma	30924	Intestinal hypomagnesemia with secondary hypocalcemia
468678	Intellectual disability-microcephaly-strabismus-behavioral abnormalities syndrome	210115	Interleukin-1 receptor antagonist deficiency	3452	Intestinal lipodystrophy
457365	Intellectual disability-muscle weakness-short stature-facial dysmorphism syndrome	169100	Interleukin-2 receptor alpha chain deficiency	3452	Intestinal lipophagic granulomatosis
3068	Intellectual disability-myopathy-short stature-endocrine defect syndrome	171208	Intermediate anorectal malformation	314376	Intestinal obstruction in the newborn due to guanylate cyclase 2C deficiency
352530	Intellectual disability-obesity-brain malformations-facial dysmorphism syndrome	268162	Intermediate BCKD deficiency	86880	Intestinal T-cell lymphoma
397973	Intellectual disability-obesity-prognathism-eye and skin anomalies syndrome	268162	Intermediate branched-chain 2-ketoacid dehydrogenase deficiency	178481	Intestinal toxemia botulism
3082	Intellectual disability-polydactyly-uncombable hair syndrome	411634	Intermediate cystinosis	178481	Intestinal toxin-mediated botulism
369837	Intellectual disability-seizures-hypotonia-ophthalmologic-skeletal anomalies syndrome	99989	Intermediate DEND syndrome	228371	Intoxication botulism
369950	Intellectual disability-seizures-macrocephaly-obesity syndrome	86797	Intermediate lichen myxedematosus	46724	Intracranial arteriovenous malformation
391372	Intellectual disability-severe speech delay-mild dysmorphism syndrome	268162	Intermediate maple syrup urine disease	252006	Intracranial endodermal sinus tumor
3409	Intellectual disability-short stature-hand contractures-genital anomalies syndrome	171433	Intermediate MSUD	91352	Intracranial germinoma
3074	Intellectual disability-short stature-hypertelorism syndrome	210110	Intermediate nemaline myopathy	252006	Intracranial yolk sac tumor
1240	Intellectual disability-short stature-wedge-shaped epiphyses of knees syndrome	309331	Intermediate osteopetrosis	137622	Intractable diarrhea-choanal atresia-eye anomalies syndrome
3051	Intellectual disability-sparse hair-brachydactyly syndrome	83418	Intermediate severe Salla disease	424058	Intraductal papillary mucinous carcinoma of pancreas
1891	Intellectual disability-spasticity-electrodactyly syndrome	279914	Intermediate spinal muscular atrophy	424982	Intrahepatic bile duct cystadenocarcinoma
		268173	Intermittent BCKD deficiency	69665	Intrahepatic cholestasis of pregnancy
		268173	Intermittent branched-chain 2-ketoacid dehydrogenase deficiency	280802	Intralobar congenital bronchopulmonary sequestration
		90283	Intermittent cutaneous lupus	280802	Intralobar congenital pulmonary sequestration
		329967	Intermittent hydrarthrosis	99088	Intramural coronary arterial course
		268173	Intermittent maple syrup urine disease	100003	Intraneural perineurioma
		268173	Intermittent MSUD	268139	Intraocular medulloepithelioma
		→2686	Intermittent neutropenia	140436	Intraosseous hemangioma
		981	Internal carotid agenesis	137686	Intrauterine adhesions
		37202	Interstitial cystitis	436144	Intrauterine growth restriction-short stature-early adult-onset diabetes syndrome
		79099	Interstitial granulomatous dermatitis with arthritis	85173	Intrauterine growth retardation-metaphyseal dysplasia-adrenal hypoplasia congenita-genital anomalies syndrome
		440427	Interstitial lung and liver disease	137686	Intrauterine synechiae
		440402	Interstitial lung disease due to ABCA3 deficiency	98839	Intravascular large B-cell lymphoma
		440402	Interstitial lung disease due to ATP-binding cassette subfamily A member 3 deficiency	98839	Intravascular lymphomatosis
				332	Intrinsic factor deficiency
				3306	Inv dup(15)

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
457088	Invasive candidiasis-deep dermatophytosis syndrome	884	Isochromosome 12p mosaicism	79143	Isolated congenital anonychia
		884	Isochromosome 12p syndrome	88620	Isolated congenital anosmia
90078	Invasive infections due to vancomycin-resistant enterococci	3307	Isochromosome 18p	162526	Isolated congenital auditory ossicle malformation
		96055	Isochromosome 21		
90078	Invasive infections due to VRE	98797	Isochromosomy Yp	180188	Isolated congenital breast hypoplasia/aplasia
99925	Invasive mole	98798	Isochromosomy Yq		
324648	Invasive non-typhoidal salmonellosis	99731	ISOD	238722	Isolated congenital controlateral synkinesia
		3306	Isodicentric 15 chromosome		
96092	Invdupdel(8p)	263524	Isolated acute necrotizing encephalopathy	217059	Isolated congenital digital clubbing
79405	Inverse JEB		99171	Isolated congenital ectropion	
329324	Inverse Klippel-Trénaunay syndrome	229717	Isolated agammaglobulinemia	432	Isolated congenital gonadotropin deficiency
		440987	Isolated agenesis of gallbladder		
98951	Inverse Marcus-Gunn phenomenon	268868	Isolated amyelia	485426	Isolated congenital hepatic fibrosis
79409	Inverse RDEB	263524	Isolated ANE	141152	Isolated congenital hypoglossia/aglossia
79409	Inverse recessive dystrophic epidermolysis bullosa	1048	Isolated anencephaly/exencephaly		
96092	Inverted 8p duplication/deletion syndrome	140989	Isolated angitis of the central nervous system	141214	Isolated congenital maxillomandibular fusion
		250923	Isolated aniridia	91489	Isolated congenital megalocornea
2704	Inverted smile-neurogenic bladder syndrome	91397	Isolated ankyloblepharon filiforme adnatum	199642	Isolated congenital microcephaly
1451	IOMID syndrome	79143	Isolated anonychia	238722	Isolated congenital mirror movements
1186	IOSCA	557	Isolated anorectal malformation	217059	Isolated congenital nail clubbing
275766	IPAH	3387	Isolated anterior cervical hypertrichosis	162516	Isolated congenital nasal pyriform aperture stenosis
747	iPAP		Isolated apertura pyriformis stenosis	91490	Isolated congenital sclerocornea
238455	IPD	162516	Isolated arhinencephaly	141214	Isolated congenital syngnathia
37042	IPEX	268936	Isolated arhinencephaly	216718	Isolated congenitally uncorrected transposition of the great arteries
88621	IPS	1166	Isolated asymmetric crying facies	216718	Isolated congenitally uncorrected transposition of the great vessels
100025	IPSID	206599	Isolated asymptomatic elevation of creatine phosphokinase	1460	Isolated CoQ-cytochrome C reductase deficiency
70592	IRAK4 deficiency	254913	Isolated ATP synthase deficiency	254905	Isolated COX deficiency
772	IRD	34528	Isolated autosomal dominant hypomagnesemia	91396	Isolated cryptophthalmia
209981	IRIDA syndrome	199326	Isolated autosomal dominant hypomagnesemia, Glaudemans type	254905	Isolated cytochrome C oxidase deficiency
64734	Iridocorneal endothelial syndrome	269221	Isolated bilateral hemispheric cerebellar hypoplasia	217	Isolated Dandy-Walker malformation
→782	Iris dysplasia-hypertelorism-deafness syndrome	158778	Isolated bone marrow mastocytosis	269212	Isolated Dandy-Walker malformation with hydrocephalus
39044	Iris melanoma	35099	Isolated brachycephaly	269215	Isolated Dandy-Walker malformation without hydrocephalus
209981	Iron-refractory iron deficiency anemia	1398	Isolated cerebellar hypoplasia/agenesis	248340	Isolated delta-SPD
43115	Iron-sulfur cluster deficiency myopathy	269203	Isolated cerebellar vermis agenesis		
86915	Irons-Bianchi syndrome	199630	Isolated cerebellar vermis hypoplasia	248340	Isolated delta-storage pool disease
209943	IRVAN syndrome	485426	Isolated CHF	248340	Isolated dense-SPD
84142	Isaac syndrome	199302	Isolated cleft lip	248340	Isolated dense-storage pool disease
84142	Isaac-Mertens syndrome	2343	Isolated cloverleaf skull syndrome	99177	Isolated distichiasis
85200	Ischio-spinal dysostosis	1460	Isolated coenzyme Q-cytochrome C reductase deficiency	35093	Isolated dolichocephaly
85200	Ischio-vertebral dysplasia	2609	Isolated complex I deficiency	1885	Isolated ectopia lentis
85200	Ischio-vertebral syndrome	217059	Isolated congenital acropachy	199647	Isolated encephalocele
1509	Ischiopatellar dysplasia	289465	Isolated congenital adermatoglyphia	221106	Isolated facial myokymia
43115	ISCU myopathy	91416	Isolated congenital alacrima	65683	Isolated focal cortical dysplasia
79144	Iso-Kikuchi syndrome	180188	Isolated congenital amastia	268994	Isolated focal cortical dysplasia type 2
79159	Isobutyric aciduria				
79159	Isobutyryl-CoA dehydrogenase deficiency				
3309	Isochromosome 5p				
3310	Isochromosome 9p				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
268961	Isolated focal cortical dysplasia type I	254913	Isolated mitochondrial respiratory chain complex V deficiency	269206	Isolated total cerebellar vermis agenesis
268973	Isolated focal cortical dysplasia type Ia	90641	Isolated mitochondrial sensorineural deafness	454750	Isolated tracheo-esophageal fistula
268980	Isolated focal cortical dysplasia type Ib	2609	Isolated NADH-coenzyme Q reductase deficiency	103909	Isolated trehalose intolerance
268987	Isolated focal cortical dysplasia type Ic	2609	Isolated NADH-CoQ reductase deficiency	238670	Isolated TRF deficiency
268994	Isolated focal cortical dysplasia type II	2609	Isolated NADH-ubiquinone reductase deficiency	238670	Isolated TRH deficiency
269001	Isolated focal cortical dysplasia type IIa	162516	Isolated nasal pyriform aperture hypoplasia	3366	Isolated trigonocephaly
269008	Isolated focal cortical dysplasia type IIb	447881	Isolated neck extensor myopathy	90674	Isolated TSH deficiency
448264	Isolated focal non-epidermolytic palmoplantar keratoderma	480556	Isolated neonatal sclerosing cholangitis	238670	Isolated TSH-releasing factor deficiency
52901	Isolated follicle stimulating hormone deficiency	137902	Isolated optic nerve hypoplasia	1460	Isolated ubiquinone-cytochrome C reductase deficiency
52901	Isolated FSH deficiency	166119	Isolated osteopoikilosis	269218	Isolated unilateral hemispheric cerebellar hypoplasia
468666	Isolated generalized anhidrosis with normal sweat glands	63440	Isolated oxycephaly	860	Isolated ventriculoarterial discordance
408	Isolated glycerol kinase deficiency	269209	Isolated partial cerebellar vermis agenesis	96	Isolated vitamin E deficiency
231662	Isolated growth hormone deficiency type IA	96269	Isolated partial vaginal agenesis	472	Isosporiasis
231671	Isolated growth hormone deficiency type IB	718	Isolated Pierre Robin sequence	2305	Isotretinoin embryopathy
231679	Isolated growth hormone deficiency type II	718	Isolated Pierre Robin syndrome	2305	Isotretinoin syndrome
231692	Isolated growth hormone deficiency type III	35098	Isolated plagiocephaly	2306	Isotretinoin-like syndrome
2128	Isolated hemihyperplasia	2924	Isolated polycystic liver disease	33	Isovaleric acid CoA dehydrogenase deficiency
2128	Isolated hemihypertrophy	2456	Isolated polythelia	33	Isovaleric acidemia
306527	Isolated hereditary congenital facial paralysis	216452	Isolated postlingual genetic deafness	309324	ISSD
229717	Isolated hypogammaglobulinemia	216445	Isolated prelingual genetic deafness	→33364	Itin syndrome
183675	Isolated IgG subclass deficiency	238670	Isolated prothyroliberin deficiency	439254	ITM2B amyloidosis
2345	Isolated Klippel-Feil syndrome	238670	Isolated protirelin deficiency	439254	ITM2B-related amyloidosis
1084	Isolated lissencephaly type 1 without known genetic defects	264691	Isolated pulmonary capillaritis	439254	ITM2B-related cerebral amyloid angiopathy
268920	Isolated macrencephaly	34528	Isolated renal magnesium wasting	435	Ito hypomelanosis
391474	Isolated median cleft face syndrome	439	Isolated right ventricular hypoplasia	3002	ITP
268920	Isolated megalencephaly	35093	Isolated scaphocephaly	457375	ITPA-related encephalopathy
238593	Isolated mesenteric lipodystrophy	440713	Isolated sedoheptulokinase deficiency	279914	IU
95707	Isolated micropenis	440713	Isolated SHPK deficiency	99123	IVC interruption
90641	Isolated mitochondrial neurosensory deafness	457083	Isolated splenogonadal fusion	294415	Ivemark II syndrome
2609	Isolated mitochondrial respiratory chain complex I deficiency	178311	Isolated sternocostoclavicular hyperostosis	97548	Ivemark syndrome
3208	Isolated mitochondrial respiratory chain complex II deficiency	3208	Isolated succinate-coenzyme Q reductase deficiency	2307	IVIC syndrome
1460	Isolated mitochondrial respiratory chain complex III deficiency	3208	Isolated succinate-CoQ reductase deficiency	281190	IWC
254905	Isolated mitochondrial respiratory chain complex IV deficiency	3208	Isolated succinate-ubiquinone reductase deficiency	3236	Jackson-Barr syndrome
		99731	Isolated sulfite oxidase deficiency	1540	Jackson-Weiss syndrome
		90674	Isolated thyroid-stimulating hormone deficiency	2848	Jacobs syndrome
		238670	Isolated thyroliberin deficiency	2308	Jacobsen syndrome
		238670	Isolated thyrotropin-releasing factor deficiency	1941	JAE
		238670	Isolated thyrotropin-releasing hormone deficiency	→636	Jaffe-Campanacci syndrome
				93277	Jaffe-Lichtenstein disease
				2269	Jagell-Holmgren-Hofer syndrome
				1873	Jalili syndrome
				300605	JALS
				73423	Jamaican vomiting sickness
				73423	Jamaican vomiting syndrome
				1891	Jancar syndrome
				2590	Jankovic-Rivera syndrome
				168491	Jansky-Bielschowsky disease

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79139	Japanese encephalitis	1454	Joubert syndrome with congenital hepatic fibrosis	199260	Juvenile aponeurotic fibromatosis
2311	Jarcho-Levin syndrome			391497	Juvenile autoimmune myasthenia gravis
474	JATD	1454	Joubert syndrome with hepatic defect	314918	Juvenile Canavan disease
91412	Jaw-winking syndrome			247794	Juvenile cataract-microcornea-renal glucosuria syndrome
313795	Jawad syndrome	397715	Joubert syndrome with JATD	300605	Juvenile Charcot disease
2315	JBS	397715	Joubert syndrome with Jeune asphyxiating thoracic dystrophy	86834	Juvenile chronic myelomonocytic leukemia
397715	JBTS with JATD			411634	Juvenile cystinosis
139431	Jeavons syndrome	220493	Joubert syndrome with ocular defect	93672	Juvenile dermatomyositis
79404	JEB-H			93672	Juvenile DM
79405	JEB-I	2318	Joubert syndrome with oculorenal defect	228254	Juvenile elastoma without osteopoikilosis
79406	JEB-lo			2929	Juvenile gastrointestinal polyposis
79402	JEB-nH gen	2754	Joubert syndrome with oral-facial-digital syndrome	98977	Juvenile glaucoma
251393	JEB-nH loc			79256	Juvenile GM1 gangliosidosis
79403	JEB-PA	2754	Joubert syndrome with orofacioidigital defect	309162	Juvenile GM2 gangliosidosis 0 variant
1201	Jejunal atresia			79230	Juvenile hemochromatosis
100077	Jejunal neuroendocrine neoplasm	220497	Joubert syndrome with renal defect	98954	Juvenile hereditary epithelial dystrophy of Meesmann
100077	Jejunal neuroendocrine tumor	220493	Joubert syndrome with retinopathy	248111	Juvenile Huntington chorea
1201	Jejunioleal atresia	2318	Joubert syndrome with Senior-Loken syndrome	248111	Juvenile Huntington disease
89840	JEN-nH			2028	Juvenile hyaline fibromatosis
→52368	Jensen syndrome	475	Joubert-Boltshauser syndrome	85435	Juvenile idiopathic rheumatoid factor-positive polyarthritis
90647	Jervell and Lange-Nielsen syndrome	2801	JPG	2929	Juvenile intestinal polyposis
33314	Jessner lymphocytic infiltration of the skin	247604	JPLS	300605	Juvenile Lou Gehrig disease
		2929	JPS	65684	Juvenile muscular atrophy of distal upper extremity
33314	Jessner-Kanof lymphocytic infiltration of the skin	2318	JS type B	65684	Juvenile muscular atrophy of the distal upper limb
		1454	JS-H	391497	Juvenile myasthenia gravis
3283	JET	220493	JS-O	86834	Juvenile myelomonocytic leukemia
		2318	JS-OR	307	Juvenile myoclonic epilepsy
474	Jeune asphyxiating thoracic dystrophy	220497	JS-R	307	Juvenile myoclonus epilepsy
		2319	Juberg-Hayward syndrome	289596	Juvenile nasopharyngeal angiofibroma
474	Jeune syndrome	101039	Juberg-Hellman syndrome	79264	Juvenile NCL
248111	JHD	93972	Juberg-Marsidi syndrome	93592	Juvenile nephronophthisis
2929	JIP	3283	Junctional ectopic tachycardia	411634	Juvenile nephropathic cystinosis
65684	JMADUE	79404	Junctional epidermolysis bullosa generalisata gravis	79264	Juvenile neuronal ceroid lipofuscinosis
307	JME			157719	Juvenile or adult CACH syndrome
86834	JMML	79402	Junctional epidermolysis bullosa generalisata mitis	85193	Juvenile osteoporosis
324999	JMP syndrome			329894	Juvenile overlap myositis
289596	JNA	79405	Junctional epidermolysis bullosa inversa	2801	Juvenile Paget disease
79264	JNCL			2801	Juvenile Paget's disease
2314	Job syndrome	79402	Junctional epidermolysis bullosa, Disentis type	247604	Juvenile PLS
2315	Johanson-Blizzard syndrome			93568	Juvenile PM
2316	Johnson neuroectodermal syndrome	79404	Junctional epidermolysis bullosa, Herlitz type	93568	Juvenile polymyositis
				79404	Junctional epidermolysis bullosa, Herlitz-Pearson type
85320	Johnson syndrome	89840	Junctional epidermolysis bullosa, non-Herlitz type	2929	Juvenile polyposis syndrome
2316	Johnson-McMillin syndrome			79403	Junctional epidermolysis bullosa-pyloric atresia syndrome
1112	Johnson-Munson syndrome	79403	Jung-Wolff-Back-Stahl syndrome	2801	Juvenile Paget disease
1485	Johnston-Aarons-Schelley syndrome			319223	Junin hemorrhagic fever
324999	Joint contractures-muscular atrophy-microcytic anemia-panniculitis-associated lipodystrophy syndrome	2321	Jung-Wolff-Back-Stahl syndrome	247604	Juvenile PLS
		989	Jussieu syndrome	93568	Juvenile PM
2295	Joint instability syndrome	1941	Juvenile absence epilepsy	93568	Juvenile polymyositis
2027	Jones syndrome	391497	Juvenile acquired myasthenia	79076	Juvenile polyposis of infancy
1256	Jorgenson-Lenz syndrome	300605	Juvenile amyotrophic lateral sclerosis	2929	Juvenile polyposis syndrome
475	Joubert syndrome				
475	Joubert syndrome type A				

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247604	Juvenile primary lateral sclerosis	2473	Kaufman-Mckusick syndrome	79141	Keratosis palmoplantaris nummularis
85436	Juvenile psoriatic arthritis	2331	Kawasaki disease	50942	Keratosis palmoplantaris striata
85408	Juvenile rheumatoid factor-negative polyarthritis	2306	Kawashima syndrome	50942	Keratosis palmoplantaris striata et areata
		2533	Kawashima-Tsuji syndrome	495	Keratosis palmoplantaris transgrediens et progrediens
247854	Juvenile rheumatoid factor-negative polyarthritis with anti-nuclear antibodies	2332	KBG syndrome	87503	Keratosis palmoplantaris transgrediens of Siemens
		439218	KCNQ2-NEE	50942	Keratosis palmoplantaris varians of Wachters
247861	Juvenile rheumatoid factor-negative polyarthritis without anti-nuclear antibodies	439218	KCNQ2-related epileptic encephalopathy	34217	Keratosis palmoplantaris with arrhythmogenic cardiomyopathy
93399	Juvenile sialidosis type 2	439218	KCNQ2-related neonatal epileptic encephalopathy	28378	Keratosis palmoplantaris-corneal dystrophy syndrome
83419	Juvenile spinal muscular atrophy	96169	KdVS	50944	Keratosis palmoplantaris-cystic eyelids-hypodontia-hypotrichosis syndrome
585	Juvenile sulfatidosis, Austin type	480	Kearns-Sayre syndrome	2198	Keratosis palmoplantaris-esophageal carcinoma syndrome
26137	Juvenile temporal arteritis	199260	Keasby tumor	2342	Keratosis palmoplantaris-periodontopathia-onychogryposis syndrome
158000	Juvenile xanthogranuloma	2662	Keipert syndrome	499	Kerion celsi
445062	Juvenile-onset diabetes mellitus-central and peripheral neurodegeneration syndrome	79233	Kelley-Seegmiller syndrome	415286	Kernicterus
		137653	Kelly-Kirson-Wyatt syndrome	3351	Kersey syndrome
79241	Juvenile-onset multiple carboxylase deficiency	54028	Kelly-Paterson syndrome	293807	Ketamine-induced biliary dilatation
		481	Kennedy disease	438075	Ketoacidosis due to monocarboxylate transporter-1 deficiency
1243	Juvenile-onset vitelliform macular dystrophy	64542	Kennedy-Teebi syndrome	1399	Ketoaciduria-intellectual disability-ataxia-deafness syndrome
		2333	Kenny syndrome	2056	Ketohexokinase deficiency
99100	Juxtaposition of the atrial appendages	2333	Kenny-Caffey syndrome	35	Ketotic hyperglycinemia
		2333	Kenny-Caffey syndrome	85202	Keutel syndrome
99100	Juxtaposition of the atrial auricles	435628	Keppen-Lubinsky syndrome	2988	Khalifa-Graham syndrome
1540	JWS	477	Keratitits-ichthyosis-deafness/Hystrix-like ichthyosis-deafness syndrome	98841	Ki-1 positive anaplastic large cell lymphoma
2322	Kabuki make-up syndrome	447777	Keratocystic odontogenic tumor	477	KID syndrome
2322	Kabuki syndrome	494	Keratoderma hereditarium mutilans	477	KID/HID syndrome
85146	Kaerer syndrome	79395	Keratoderma hereditarium mutilans with ichthyosis	97332	Kienbock disease
29073	Kahler's disease	34217	Keratoderma with woolly hair type I	50918	Kikuchi disease
→324737	Kahrizi syndrome	65282	Keratoderma with woolly hair type II	50918	Kikuchi-Fujimoto disease
2324	Kaler-Garrity-Stern syndrome	420686	Keratoderma with woolly hair type IV	482	Kimura disease
2325	Kallin syndrome	79395	Keratoderma-ichthyosiform dermatosis-elevated beta-glucuronidase syndrome	401996	KIN
478	Kallmann syndrome	79501	Keratoderma palmoplantaris papulosa, Buschke-Fischer-Brauer type	2908	Kindler syndrome
2326	Kallmann syndrome-heart disease syndrome	50943	Keratolytic winter erythema	99741	King-Denborough syndrome
99179	Kandori fleck retina	495	Keratosis extremitatum hereditaria progrediens	565	Kinky hair disease
1836	Kantaputra mesomelic dysplasia	218	Keratosis follicularis	565	Kinky hair syndrome
79280	Kanzaki disease	2340	Keratosis follicularis spinulosa decalvans	1183	Kinsbourne syndrome
949	Kaplan-Plauchu-Fitch syndrome	2339	Keratosis follicularis-dwarfism-cerebral atrophy syndrome	100996	Kjellin syndrome
→3157	Kaplowitz-Bodurtha syndrome	281201	Keratosis linearis-ichthyosis congenita-sclerosing keratoderma syndrome	98673	Kjer optic atrophy
33276	Kaposi sarcoma	86919	Keratosis palmaris et plantaris-clinodactyly syndrome	99978	Klatskin tumor
2122	Kaposiform hemangioendothelioma	678	Keratosis palmoplantar-periodontopathy syndrome	261494	Kleefstra syndrome
464329	Kaposiform lymphangiomatosis				
183675	Kappa-chain deficiency				
2328	Kapur-Toriello syndrome				
1381	Karandikar-Maria-Kamble syndrome				
2329	Karsch-Neugebauer syndrome				
→244	Kartagener syndrome				
401996	Karyomegalic interstitial nephritis				
2330	Kasabach-Merritt syndrome				
1894	Kasznica-Carlson-Coppedge syndrome				
3360	Katsantoni-Papadakou Lagoyanni syndrome				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
96147	Kleefstra syndrome due to 9q subtelomeric deletion	629	Kowarski syndrome	2363	Lacrimoauriculodentodigital syndrome
96147	Kleefstra syndrome due to 9q34 microdeletion	2352	Kozlowski-Brown-Hardwick syndrome	2363	Lacrimoauricularodigital syndrome
261652	Kleefstra syndrome due to a point mutation	3082	Kozlowski-Krajewska syndrome	284426	Lactate dehydrogenase A deficiency
96147	Kleefstra syndrome due to del(9)(q34)	2204	Kozlowski-Tsuruta syndrome	284435	Lactate dehydrogenase B deficiency
96147	Kleefstra syndrome due to monosomy 9q34	487	Krabbe disease	2965	Lactotroph adenoma
896	Klein-Waardenburg syndrome	206436	Krabbe disease, classic form	2968	LAD
33543	Kleine-Levin syndrome	206436	Krabbe disease, early-onset	99844	LAD-1 variant
2110	Kleiner-Holmes syndrome	206443	Krabbe disease, late-onset	99842	LAD-I
399081	KLHL9-related early-onset distal myopathy	1345	Krasnow-Qazi syndrome	99843	LAD-II
281201	KLICK syndrome	709	Krause-Kivlin syndrome	99844	LAD-III
447974	Klippel-Feil anomaly-myopathy-facial dysmorphism syndrome	709	Krause-van Schooneveld-Kivlin syndrome	2363	LADD syndrome
2345	Klippel-Feil malformation	284149	Kreiborg-Pakistani syndrome	1484	Ladda-Zonana-Ramer syndrome
2345	Klippel-Feil sequence	89838	KRT14-related epidermolysis bullosa simplex	158687	LAEB
90308	Klippel-Trénaunay syndrome	2908	KS	501	Lafora disease
2346	Klippel-Trénaunay-Weber syndrome	293936	KTCNCT	1997	Lagophthalmia-cleft lip and palate syndrome
157823	Klüver-Bucy syndrome	447777	KTOC	59135	Laing early-onset distal myopathy
485	Kniest dysplasia	306674	Kufor-Rakeb syndrome	275761	LAL deficiency
1571	Knobloch syndrome	79262	Kufs disease	538	LAM
1571	Knobloch-Layer syndrome	83419	Kugelberg-Welander disease	306507	LAMB2-related infantile-onset nephrotic syndrome
2698	Knuckle pads-leukonychia-sensorineural deafness-palmoplantar hyperkeratosis syndrome	→1487	Kumar-Levick syndrome	1296	Lambert syndrome
2698	Knuckle pads-leukonychia-sensorineural deafness-palmoplantar keratoderma syndrome	2505	Kunze-Riehm syndrome	43393	Lambert-Eaton myasthenic syndrome
2349	Kocher-Debré-Semelaigne syndrome	→794	Kurczynski-Casperson syndrome	313	Lamellar ichthyosis
679	Köhlmeier-Degos disease	454745	Kuru	137871	Laminopathy type Decaudain-Vigouroux
679	Köhlmeier-Degos-Delort-Tricort syndrome	1149	Kuskokwim disease	137871	Laminopathy with severe metabolic syndrome and myopathy
1946	Kohlschütter-Tonz syndrome	767	Küssmaul-Maier disease	90024	LAMM syndrome
3197	Kok disease	449432	Küttner tumor	98818	Landau-Kleffner syndrome
99077	Kommerell diverticulum	2798	Kuzniecky syndrome	354	Landing disease
3130	Komuragaeri disease	34217	KWWH type I	269	Landouzy-Dejerine myopathy
2764	König disease	65282	KWWH type II	231031	Lane disease
→1215	Königsmark-Knox-Hussels syndrome	420686	KWWH type IV	2632	Langer mesomelic dysplasia
96169	Koolen-De Vries syndrome	319254	Kyasanur forest disease	502	Langer-Giedion syndrome
363965	Koolen-De Vries syndrome due to a point mutation	319254	Kyasanur hemorrhagic fever	86897	Langerhans cell sarcoma
2892	Kopysc-Barczyk-Krol syndrome	79155	Kynureninase deficiency	2368	Laparoschisis
477831	Kosaki overgrowth syndrome	1801	Kyphomelic dysplasia	→1159	Laplane-Fontaine-Lagardere syndrome
2839	Kosenow syndrome	275543	L1 syndrome	2363	LARD syndrome
99749	Kostmann syndrome	275543	L1CAM syndrome	98838	Large cell lymphoma of the mediastinum
1129	Kosztolanyi syndrome	79314	L-2-HGA	626	Large congenital melanocytic nevus
99741	Koussef-Nichols syndrome	79314	L-2-hydroxyglutaric acidemia	633	Laron syndrome
2351	Kousseff syndrome	79314	L-2-hydroxyglutaric aciduria	220465	Laron syndrome with immunodeficiency
		35704	L-Arginine:glycine amidinotransferase deficiency	220465	Laron-like syndrome
		157973	L-CMD	633	Laron-type dwarfism
		156	L-CPT1 deficiency	2370	Larsen-like osseous dysplasia-short stature syndrome
		156	L-CPTI deficiency	284139	Larsen-like syndrome, B3GAT3 type
		440731	L-ferritin deficiency		
		93599	L-glyceric aciduria		
		216694	L-transposition of the great arteries		
		83483	La Crosse encephalitis		
		53696	LAABHD		
		3473	Laband syndrome		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2808	Laryngeal abductor paralysis	98912	Late-onset distal myopathy, Markesbery-Griggs type	104	Leber optic atrophy
2375	Laryngeal abductor paralysis-intellectual disability syndrome	228227	Late-onset focal dermal elastosis	99718	Leber plus disease
2407	Laryngeal and ocular granulation tissue in children from the Indian subcontinent syndrome	163708	Late-onset infantile spasms	98955	LECD
		199299	Late-onset isolated ACTH deficiency	650	Lecithin-cholesterol acyltransferase deficiency
100083	Laryngeal neuroendocrine tumor	79406	Late-onset junctional epidermolysis bullosa	199251	Ledderhose disease
2407	Laryngo-onycho-cutaneous syndrome	231556	Late-onset localized junctional epidermolysis bullosa-intellectual disability syndrome	71273	Left renal vein entrapment syndrome
2004	Laryngo-tracheo-esophageal cleft			99111	Left superior caval vein persisting to left-sided atrium
280205	Laryngo-tracheo-esophageal cleft type 0	79241	Late-onset multiple carboxylase deficiency	99111	Left superior vena cava persisting to left-sided atrium
93938	Laryngo-tracheo-esophageal cleft type 1	93589	Late-onset nephronophthisis	99111	Left SVC persisting to left-sided atrium
		90186	Late-onset primary lymphedema		
93939	Laryngo-tracheo-esophageal cleft type 2	67042	Late-onset retinal degeneration	54260	Left ventricular hypertrabeculation
		2789	Lateral meningocele syndrome	54260	Left ventricular noncompaction
93940	Laryngo-tracheo-esophageal cleft type 3	141136	Laterofacial microsomia	99095	Left ventricular-to-right atrial communication
		46059	Lathosterolosis		
93941	Laryngo-tracheo-esophageal cleft type 4	98964	Lattice corneal dystrophy type 1	1757	Leg duplication-mirror foot syndrome
		98964	Lattice corneal dystrophy type I		
2005	Laryngo-tracheo-esophageal cleft-pulmonary hypoplasia syndrome	99094	Laubry-Pezzi syndrome	2380	Legg-Calvé-Perthes disease
2004	Laryngo-tracheo-esophageal diastema	2398	Launois-Bensaude lipomatosis	549	Legionellosis
		2377	Laurence-Moon syndrome	549	Legionnaires disease
2372	Laryngocele	2378	Laurin-Sandrow syndrome	137605	Legius syndrome
137935	Laryngotracheal angioma	79086	Lawrence syndrome	2789	Lehman syndrome
2004	Laryngotracheoesophageal cleft	79086	Lawrence-Seip syndrome	1647	Leichtman-Wood-Rohn syndrome
280205	Laryngotracheoesophageal cleft type 0	2379	Laxova-Opitz syndrome	255241	Leigh disease with leukodystrophy
		137898	LBSL	70474	Leigh disease with myopathy
93938	Laryngotracheoesophageal cleft type 1	2369	LBWC syndrome	255249	Leigh disease with nephrotic syndrome
		2004	LC		
93939	Laryngotracheoesophageal cleft type 2	99900	LCAD	3008	Leigh necrotizing encephalopathy due to pyruvate carboxylase deficiency
		650	LCAT deficiency		
93940	Laryngotracheoesophageal cleft type 3	1486	LCCS1	3008	Leigh syndrome due to PC deficiency
		137776	LCCS2		
93941	Laryngotracheoesophageal cleft type 4	137783	LCCS3	3008	Leigh syndrome due to pyruvate carboxylase deficiency
		98964	LCD1		
1202	Larynx atresia	93558	LCDD	70474	Leigh syndrome with cardiomyopathy
99824	Lassa fever	98964	LCDI		
99824	Lassa hemorrhagic fever	5	LCHAD deficiency	255241	Leigh syndrome with leukodystrophy
98974	Late hereditary endothelial dystrophy	5	LCHADD		
		52416	LCM	255249	Leigh syndrome with nephrotic syndrome
157716	Late infantile CACH syndrome	626	LCMN		
168491	Late infantile NCL	363618	LCPS	70472	Leigh syndrome, French-Canadian type
168491	Late infantile neuronal ceroid lipofuscinosis	65285	LDD		
		2364	LDH deficiency	70472	Leigh syndrome, Saguenay-Lac-Saint-Jean type
79256	Late-infantile GM1 gangliosidosis	284435	LDH-H subunit deficiency		
206443	Late-infantile/juvenile Krabbe disease	284426	LDH-M subunit deficiency	485421	Leigh-like basal ganglia disease-optic atrophy-peripheral neuropathy syndrome
		2616	Le Merrer syndrome		
98816	Late-onset benign childhood occipital epilepsy	330015	Lead intoxication	314	Leiner disease
		330015	Lead poisoning		
247573	Late-onset citrullinemia type 1	3246	Learman syndrome	71274	Leiomyomatosis peritonealis disseminate
247573	Late-onset citrullinemia type I	65	Leber congenital amaurosis	64720	Leiomyosarcoma
399058	Late-onset distal crystallinopathy	104	Leber hereditary optic neuropathy		
		190	Leber miliary aneurysm	104076	Leiomyosarcoma of small intestine

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
213807	Leiomyosarcoma of the cervix uteri	1237	Lethal hydrocephalus-cardiac malformation-dense bones syndrome	439224	Leukocyte chemotactic factor-2 amyloidosis
213625	Leiomyosarcoma of the corpus uteri			77295	Leukodystrophy with oligodontia
507	Leishmaniasis	35064	Lethal idiopathic viral infection	139444	Leukoencephalopathy with bilateral anterior temporal lobe cysts
140936	Leleis syndrome				
137839	Lemierre postanginal sepsis	254857	Lethal infantile mitochondrial disease	137898	Leukoencephalopathy with brain stem and spinal cord involvement-high lactate syndrome
137839	Lemierre syndrome				
2382	Lennox-Gastaut syndrome	254857	Lethal infantile mitochondrial myopathy	137898	Leukoencephalopathy with brain stem and spinal cord involvement-lactate elevation syndrome
209959	Lens-induced endophthalmitis				
209959	Lens-induced iridocyclitis	2347	Lethal Kniest-like dysplasia	137898	Leukoencephalopathy with mild cerebellar ataxia and white matter edema
209959	Lens-induced uveitis				
568	Lenz microphthalmia	478049	Lethal left ventricular non-compaction-seizures-hypotonia-cataract-developmental delay syndrome	363540	Leukoencephalopathy with vanishing white matter
2658	Lenz-Majewski hyperostotic dwarfism				
500	LEOPARD syndrome	86879	Lethal midline granuloma	135	Leukoencephalopathy with vanishing white matter
330032	Lepore-beta-thalassemia syndrome				
508	Leprechaunism	33108	Lethal multiple pterygium syndrome	137639	Leukoencephalopathy-ataxia-hypodontia-hypomyelination syndrome
548	Leprosy				
252031	Leptomeningeal melanomatosis	435845	Lethal neonatal rigidity-multifocal seizure syndrome	163684	Leukoencephalopathy-dystonia-motor neuropathy syndrome
268838	Leptomylolipoma				
509	Leptospirosis	300313	Lethal neurodegenerative disorder due to copper transport defect	83629	Leukoencephalopathy-metaphyseal chondrodysplasia syndrome
2900	Leri pleonosteosis				
240	Léri-Weill dyschondrosteosis	293925	Lethal occipital encephalocele-skeletal dysplasia syndrome	2386	Leukoencephalopathy-palmo-plantar keratoderma syndrome
240	Léri-Weill syndrome				
510	Lesch-Nyhan syndrome	2736	Lethal omphalocele-cleft palate syndrome	314051	Leukoencephalopathy-thalamus and brainstem anomalies-high lactate syndrome
158687	Lethal acantholytic epidermolysis bullosa				
314718	Lethal arteriopathy syndrome due to fibulin-4 deficiency	216804	Lethal osteogenesis imperfecta	1816	Leukomelanoderma-intellectual disability-hypotrichosis syndrome
53696	Lethal arthrogyposis-anterior horn cell disease syndrome				
1187	Lethal ataxia with deafness and optic atrophy	1832	Lethal osteosclerotic bone dysplasia	2387	Leukonychia totalis
1420	Lethal chondrodysplasia, Moerman type				
1421	Lethal chondrodysplasia, Seller type	210144	Lethal polymalformative syndrome, Boissel type	210133	Leukonychia totalis-acanthosis-nigricans-like lesions-abnormal hair syndrome
1486	Lethal congenital contracture syndrome type 1				
137776	Lethal congenital contracture syndrome type 2	1234	Lethal popliteal pterygium syndrome	2045	Leukonychia totalis-trichilemmal cysts-ciliary dystrophy syndrome
137783	Lethal congenital contracture syndrome type 3				
330050	Lethal encephalopathy due to mitochondrial and peroxisomal fission defect	1423	Lethal recessive chondrodysplasia	79507	Leukotriene C4 synthase deficiency
1972	Lethal faciocardiomelic dysplasia				
444069	Lethal fetal brain malformation-duodenal atresia-bilateral renal hypoplasia syndrome	1662	Lethal restrictive dermopathy	2743	Levic-Stefanovic-Nikolic syndrome
439897	Lethal fetal cerebrogenitourinary agenesis/hypoplasia syndrome				
1046	Lethal hemolytic anemia-genital anomalies syndrome	→56304	Lethal short-limb dwarfism, McAlister-Crane type	2388	Levine-Critchley syndrome
480528	Lethal hydranencephaly-diaphragmatic hernia syndrome				
		464366	Lethal skeletal dysplasia-fetal akinesia-contractures-thoracic dysplasia-pulmonary hypoplasia syndrome	216694	Levo-transposition of the great arteries
		79022	Lethal variant of Simpson-Golabi-Behmel syndrome	95854	Levocardia
		99870	Letterer-Siwe disease	95854	Levocardia-situs inversus
		58017	Leukemic reticuloendotheliosis	2363	Levy-Hollister syndrome
		300878	Leukemic reticuloendotheliosis variant	302	Lewandowsky-Lutz syndrome
		2968	Leukocyte adhesion deficiency	→1896	Lewis-Pashayan syndrome
		99842	Leukocyte adhesion deficiency type I	48162	Lewis-Sumner syndrome
		99843	Leukocyte adhesion deficiency type II	755	Leydig cell hypoplasia
		99844	Leukocyte adhesion deficiency type III	96265	Leydig cell hypoplasia due to complete LH receptor inactivation
		99844	Leukocyte adhesion deficiency-1 variant	96265	Leydig cell hypoplasia due to complete LH resistance
				96265	Leydig cell hypoplasia due to complete luteinizing hormone receptor inactivation

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96265	Leydig cell hypoplasia due to complete luteinizing hormone resistance	104	LHON	34514	Limb-girdle muscular dystrophy due to telethonin deficiency
		99718	LHON plus disease		
		313	LI		
325448	Leydig cell hypoplasia due to LHB deficiency	524	Li-Fraumeni syndrome	1878	Limb-girdle muscular dystrophy due to TRIM32 deficiency
		49804	Lichen amyloidosis	257	Limb-girdle muscular dystrophy with epidermolysis bullosa simplex
325448	Leydig cell hypoplasia due to luteinizing hormone subunit beta deficiency	49804	Lichen amyloidosis		
		525	Lichen follicularis		
		525	Lichen planopilaris		
96266	Leydig cell hypoplasia due to partial LH receptor inactivation	254395	Lichen planus actinus	86812	Limb-girdle muscular dystrophy-intellectual disability syndrome
96266	Leydig cell hypoplasia due to partial LH resistance	525	Lichen planus follicularis	69085	Limb-mammary syndrome
		254478	Lichen planus pemphigoides	171673	Limb stem cell deficiency
96266	Leydig cell hypoplasia due to partial luteinizing hormone receptor inactivation	254463	Lichen planus pigmentosa	276402	Limbic encephalitis with caspr2 antibodies
		254463	Lichen planus pigmentosus		
		254463	Lichen planus pigmentosus inversus		
		254395	Lichen planus subtropicus		
96266	Leydig cell hypoplasia due to partial luteinizing hormone resistance	254395	Lichen planus tropicus	329341	Limbic encephalitis with dipeptidyl-peptidase 6 antibodies
		254395	Lichenoid melanodermitis	329341	Limbic encephalitis with DPP6 antibodies
99824	LF	2390	Lichtenstein syndrome	329341	Limbic encephalitis with DPPX antibodies
266	LGMD1A	448251	Lichtenstein-Knorr syndrome		
264	LGMD1B	526	Liddle syndrome	163908	Limbic encephalitis with leucine-rich glioma-inactivated 1 antibodies
265	LGMD1C	1275	Liebenberg syndrome		
34516	LGMD1D	99812	LIG4 syndrome	163908	Limbic encephalitis with LGI1 antibodies
34517	LGMD1E	99812	Ligase 4 syndrome		
55595	LGMD1F	93557	Light and heavy chain deposition disease	217253	Limbic encephalitis with N-methyl-D-aspartate receptor antibodies
55596	LGMD1G				
238755	LGMD1H	93558	Light chain deposition disease	163914	Limbic encephalitis with nCMAgs antibodies
267	LGMD2A	85443	Light-chain amyloidosis		
268	LGMD2B	97231	Ligneous conjunctivitis	217253	Limbic encephalitis with NMDA receptor antibodies
353	LGMD2C	2369	Limb body wall complex		
62	LGMD2D	2492	Limb transversal defect-cardiac anomaly syndrome	163914	Limbic encephalitis with novel cell membrane antigen antibodies
119	LGMD2E				
219	LGMD2F	974	Limb, scalp and skull defects	83467	Limbic encephalitis-neuromyotonia-hyperhidrosis-polyneuropathy syndrome
34514	LGMD2G	62	Limb-girdle muscular dystrophy due to alpha-sarcoglycan deficiency		
1878	LGMD2H	119	Limb-girdle muscular dystrophy due to beta-sarcoglycan deficiency	254857	LIMD
34515	LGMD2I			366	Limit dextrinosis
140922	LGMD2J	267	Limb-girdle muscular dystrophy due to calpain deficiency	220402	Limited cutaneous systemic sclerosis
86812	LGMD2K			265	Limb-girdle muscular dystrophy due to caveolin-3 deficiency
206549	LGMD2L	219	Limb-girdle muscular dystrophy due to delta-sarcoglycan deficiency		
206554	LGMD2M			268	Limb-girdle muscular dystrophy due to dysferlin deficiency
206559	LGMD2N	34515	Limb-girdle muscular dystrophy due to FKRP deficiency		
206564	LGMD2O			79150	Limb-girdle muscular dystrophy due to gamma-sarcoglycan deficiency
280333	LGMD2P	353	Limb-girdle muscular dystrophy due to lamin A/C deficiency		
254361	LGMD2Q			264	Limb-girdle muscular dystrophy due to myotilin deficiency
363543	LGMD2R	266	Limb-girdle muscular dystrophy due to POMK deficiency		
369840	LGMD2S			445110	Limb-girdle muscular dystrophy due to POMK deficiency
363623	LGMD2T	2612	Limb-girdle muscular dystrophy due to POMK deficiency		
352479	LGMD2U			2611	Limb-girdle muscular dystrophy due to POMK deficiency
466801	LGMD2W	2611	Limb-girdle muscular dystrophy due to POMK deficiency		
476084	LGMD2X			2611	Limb-girdle muscular dystrophy due to POMK deficiency
424261	LGMD2Y	2611	Limb-girdle muscular dystrophy due to POMK deficiency		
480682	LGMD2Z				
445110	LGMD due to POMK deficiency				
93557	LHCDD				
65285	Lhermitte-Duclos disease				

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36273	Linitis plastica of the stomach	100011	Lissencephaly with cerebellar hypoplasia type A	2407	LOGIC syndrome
888	Lip-pit syndrome			250831	Logopenic primary progressive aphasia
435660	LIPE-related familial partial lipodystrophy	100012	Lissencephaly with cerebellar hypoplasia type B	250831	Logopenic progressive aphasia
435660	LIPE-related FPLD	100013	Lissencephaly with cerebellar hypoplasia type C	250831	Logopenic variant PPA
77243	Lipedema			2404	Loiasis
255182	Lipoamide dehydrogenase deficiency	100014	Lissencephaly with cerebellar hypoplasia type D	5	Long chain 3-hydroxyacyl-CoA dehydrogenase deficiency
528	Lipoatrophic diabetes	100015	Lissencephaly with cerebellar hypoplasia type E	99900	Long chain acyl-CoA dehydrogenase deficiency
156156	Lipoatrophy with diabetes, leukomelanodermic papules, liver steatosis, and hypertrophic cardiomyopathy	100016	Lissencephaly with cerebellar hypoplasia type F	3363	Long eyelashes-intellectual disability syndrome
		533	Listeriosis	90647	Long QT interval-deafness syndrome
247762	Lipoblastoma	820	Livedo racemosa-cerebrovascular accident syndrome	37553	Long QT syndrome type 7
90156	Lipodystrophia centrifugalis abdominalis infantilis	820	Livedo reticularis-cerebrovascular accident syndrome	65283	Long QT syndrome type 8
1979	Lipodystrophy due to peptidic growth factors deficiency	79095	Liver disease-retinitis pigmentosa-polyneuropathy-epilepsy syndrome	65283	Long QT syndrome-syndactyly syndrome
50811	Lipodystrophy-intellectual disability-deafness syndrome	369	Liver glycogen phosphorylase deficiency	5	Long-chain 3-hydroxyacyl-coenzyme A dehydrogenase deficiency
3163	Lipodystrophy-Rieger anomaly-diabetes syndrome	98818	LKS	180157	Longitudinal vaginal septum
401859	Lipoic acid synthetase deficiency	363618	LMNA-related cardiocutaneous progeria syndrome	52054	Longman-Tolmie syndrome
139436	Lipoid dermatoarthritis			168	Loose anagen syndrome
530	Lipoid proteinosis	157973	LMNA-related congenital muscular dystrophy	411602	LOPD
36397	Lipomatosis dolorosa	33108	LMPS	2832	Lopes-Gorlin syndrome
238593	Lipomatous mesenteritis	69085	LMS	2266	Lopes-Marques de Faria syndrome
812	Lipomucopolysaccharidosis	93924	Lobar holoprosencephaly	67042	LORD
268835	Lipomyelomeningocele	666	Lobstein disease	79395	Loricrin keratoderma
329481	Lipoprotein glomerulopathy	2440	Lobster-claw deformity	803	Lou Gehrig disease
69078	Liposarcoma	2407	LOC syndrome	100	Louis-Bar syndrome
238593	Liposclerotic mesenteritis	314709	Localized AL amyloidosis	171215	Low anorectal malformation
401862	Lipoyl transferase 1 deficiency	93685	Localized Castleman disease	251633	Low grade ependymoma
447795	Lipoyl transferase 2 deficiency	263534	Localized deciduous skin	69663	Low phospholipid associated cholelithiasis
98955	Lisch epithelial corneal dystrophy	79400	Localized epidermolysis bullosa simplex	140949	Low-flow priapism
2400	Lisker-Garcia-Ramos syndrome	90289	Localized fibrosing scleroderma	213736	Low-grade neuroendocrine tumor of the corpus uteri
101003	Lison syndrome	251393	Localized junctional epidermolysis bullosa, non-Herlitz type	213736	Low-grade neuroendocrine tumor of the uterine corpus
531	Lissencephaly due to 17p13.3 deletion	90398	Localized lichen myxedematosus with mixed features of different subtypes	1652	Low-molecular-weight proteinuria with hypercalciuria and nephrocalcinosis
95232	Lissencephaly due to LIS1 mutation	90399	Localized lichen myxedematosus with monoclonal gammopathy or systemic symptoms	534	Low disease
171680	Lissencephaly due to TUBA1A mutation			534	Low eye disease
89844	Lissencephaly syndrome, Norman-Roberts type	178517	Localized pagetoid reticulosis	534	Low eye disease
2148	Lissencephaly type 1 due to doublecortin gene mutation	263534	Localized PSS	2408	Low eye disease
352682	Lissencephaly type 2 without muscular or eye involvement	163927	Localized pustular psoriasis	363447	Lower extremity-predominant autosomal dominant proximal spinal muscular atrophy
		90289	Localized scleroderma		
352682	Lissencephaly type 2 without muscular or ocular involvement	2406	Locked-in syndrome	363454	Lower extremity-predominant autosomal dominant proximal spinal muscular atrophy with contractures
86821	Lissencephaly type 3-familial fetal akinesia sequence syndrome	75566	Loeffler endocarditis		
		724	Loeffler syndrome		
86822	Lissencephaly type 3-metacarpal bone dysplasia syndrome	60030	Loeys-Dietz syndrome		
		724	Loeffler syndrome		

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209341	Lower extremity-predominant autosomal dominant proximal spinal muscular atrophy without contractures	1120	Lung agenesis-heart defect-thumb anomalies syndrome	280302	Lymphoplasmacytic sclerosing pancreatitis
		137631	Lung fibrosis-immunodeficiency-46,XX gonadal dysgenesis syndrome	144	Lynch syndrome
2487	Lower limb deficiency-hypospadias syndrome	90285	Lupus erythematosus panniculitis	1123	Lynch-Lee-Murday syndrome
		90285	Lupus erythematosus profundus	3196	Lyngstadaas syndrome
295051	Lower limb hypertrophy	90283	Lupus erythematosus tumidus	98842	LyP
141064	Lower lip fistula	1173	Luteinizing hormone-releasing hormone deficiency with ataxia	2203	Lysine alpha-ketoglutarate reductase deficiency
276435	Lower motor neuron syndrome with late-adult onset		Lutz-Lewandowsky epidermodysplasia verruciformis	470	Lysinuric protein intolerance
844	Lown-Ganong-Levine syndrome	302	Lutz-Richner-Landolt syndrome	275761	Lysosomal acid lipase deficiency
1533	Lowry syndrome	→2697	LVNC	61	Lysosomal alpha-D-mannosidase deficiency
2409	Lowry-MacLean syndrome	54260	Lyell syndrome	309288	Lysosomal alpha-D-mannosidase deficiency, adult form
1824	Lowry-Wood syndrome	537	LYG	309282	Lysosomal alpha-D-mannosidase deficiency, infantile form
2003	Lowry-Yong syndrome	86869	Lyme borreliosis	34587	Lysosomal glycogen storage disease with normal acid maltase activity
254478	LP pemphigoides	91546	Lyme disease		
254463	LP pigmentosa	91546	Lymphangioliomyomatosis	79284	Lysosomal membrane cobalamin transporter deficiency
254463	LP pigmentosus	538	Lymphatic filariasis	93561	Lysozyme amyloidosis
250831	LPA	2035	Lymphedema praecox	90020	Lytic-Bodig disease
69663	LPAC	→289825	Lymphedema tarda	330041	M hemoglobinopathy
71274	LPD	→289825	Lymphedema with yellow nails	247262	Mabry syndrome
329481	LPG	662	Lymphedema-atrial septal defects-facial changes syndrome	98938	MAC
470	LPI	86915	Lymphedema-cerebral arteriovenous anomaly syndrome	36412	Mac Duffie hypocomplementemic urticarial vasculitis
309015	LPL deficiency	33001	Lymphedema-distichiasis syndrome	36412	Mac Duffie syndrome
163927	LPP	1563	Lymphedema-hypparathyroidism syndrome syndrome	2220	MacDermot-Patton-Williams syndrome
525	LPP		Lymphedema-lymphangiectasia-intellectual disability syndrome	2083	MacDermot-Winter syndrome
37553	LQT7	2136	Lymphedema-posterior choanal atresia syndrome	98757	Machado disease
65283	LQT8	99141	Lymphedema-ptosis syndrome	98757	Machado-Joseph disease
314051	LTBL	→33001	Lymphoadenopathic mastocytosis with eosinophilia	276238	Machado-Joseph disease type 1
79507	LTC4 synthase deficiency	158793	Lymphoblastoid variant of NK-cell lymphoma	276241	Machado-Joseph disease type 2
2004	LTEC		Lymphocytic hypereosinophilic syndrome	276244	Machado-Joseph disease type 3
280205	LTEC0	86870	Lymphocytic interstitial pneumonia	319229	Machupo hemorrhagic fever
93938	LTEC1	314970	Lymphocytic variant HES	79495	Macias Flores-Garcia Cruz-Rivera syndrome
93939	LTEC2	79128	Lymphoepithelial-like carcinoma	1574	Mackay-Shek-Carr syndrome
93940	LTEC3	314970	Lymphogranulomatosis X	468672	MACOM syndrome
93941	LTEC4	289682	Lymphoid HES	357158	Macroblepharon-ectropion-hypertelorism-macrostromia syndrome
93938	LTEC I	86886	Lymphoid interstitial pneumonia	217335	Macrocephaly-alopecia-cutis laxa-scoliosis syndrome
93939	LTEC II	79128	Lymphomatoid granulomatosis	60040	Macrocephaly-capillary malformation syndrome
93940	LTEC III	86869	Lymphomatoid papulosis		
93941	LTEC IV	98842	Lymphomatous meningitis	60040	Macrocephaly-cutis marmorata telangiectatica congenita syndrome
53351	Lubag	329998	Lymphoplasmacytic lymphoma without IgM production	397612	Macrocephaly-developmental delay syndrome
53351	Lubag syndrome	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
2575	Lubani-Al Saleh-Teebi syndrome	443159	Lymphoplasmacytic lymphoma without IgM production	210548	Macrocephaly-intellectual disability-autism syndrome
2410	Lubinsky syndrome				
→1762	Lubs-Arena syndrome	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
2312	Lucey-Driscoll syndrome	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
776	Lujan syndrome	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
776	Lujan-Fryns syndrome	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
319213	Lujo hemorrhagic fever	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
268388	Lumbosacral spina bifida aperta	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
268758	Lumbosacral spina bifida cystica	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
97332	Lunatomalacia	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome
2928	Lundberg syndrome	443159	Lymphoplasmacytic lymphoma without Immunoglobulin M production	210548	Macrocephaly-intellectual disability-autism syndrome

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
466791	Macrocephaly-intellectual disability-left ventricular non compaction syndrome	394532	MADD, mild type	755	Male pseudohermaphroditism due to luteinizing hormone resistance or luteinizing hormone beta subunit deficiency
		394529	MADD, severe neonatal type		
		35688	Madelung deformity		
457485	Macrocephaly-intellectual disability-neurodevelopmental disorder-small thorax syndrome	295223	Madelung deformity, bilateral	1646	Male sterility due to chromosome Y deletion
		295221	Madelung deformity, unilateral		
		2398	Madelung disease		
2563	Macrocephaly-obesity-mental disability-ocular abnormalities syndrome	137867	Madras motor neuron disease	3000	Male-limited precocious puberty
		48162	MADSAM	99915	Malignant granulosa cell tumor of ovary
		2583	Madura foot	289385	Malignancy diagnosed during pregnancy
		1942	MAE		
2427	Macrocephaly-short stature-paraplegia syndrome	171709	Mae infertility due to round-headed spermatozoa	98839	Malignant angioendotheliomatosis
2429	Macrocephaly-spastic paraplegia-dysmorphism syndrome	199354	Maeda syndrome	679	Malignant atrophic papulosis
79489	Macrocytic lymphangioma	163634	Maffucci syndrome	100093	Malignant carcinoid syndrome
79489	Macrocytic lymphatic malformation	324972	MAGIC syndrome	99912	Malignant dysgerminomatous germ cell tumor of ovary
295044	Macroductyly of fingers	438274	Mahvash disease	276145	Malignant epithelial tumor of salivary glands
295241	Macroductyly of fingers, bilateral	77297	Majeed syndrome		
295239	Macroductyly of fingers, unilateral	2637	Majewski osteodysplastic primordial dwarfism type II	213837	Malignant germ cell tumor of cervix uteri
295047	Macroductyly of foot	70470	Major hyperlipidemia	213751	Malignant germ cell tumor of corpus uteri
295245	Macroductyly of foot, bilateral	210272	Mal de débarquement		
295243	Macroductyly of foot, unilateral	87503	Mal de Meleda	423	Malignant hyperthermia of anesthesia
295044	Macroductyly of hand	556	Malakoplakia		
295241	Macroductyly of hand, bilateral	420179	Malan overgrowth syndrome	2215	Malignant hyperthermia-arthrogryposis-torticollis syndrome
295239	Macroductyly of hand, unilateral	673	Malaria		
295047	Macroductyly of toes	75376	Malattia leventinese	168999	Malignant melanoma of the mucosa
295245	Macroductyly of toes, bilateral	401973	Male EBP disorder with neurological defects	293181	Malignant migrating partial epilepsy of infancy
295243	Macroductyly of toes, unilateral			293181	Malignant migrating partial seizures of infancy
2477	Macroencephaly	2234	Male hypergonadotropic hypogonadism-intellectual disability-skeletal anomalies syndrome	213610	Malignant mixed Müllerian tumor of the corpus uteri
158061	Macrophage activation syndrome			213512	Malignant mixed Müllerian tumor of the ovary
592	Macrophagic myofasciitis	171709	Male infertility due to globozoospermia	213787	Malignant Müllerian mixed tumor of the cervix uteri
2432	Macrosomia-microphthalmia-cleft palate syndrome	137893	Male infertility due to large-headed multiflagellar polyploid spermatozoa	3148	Malignant neurilemmoma
2563	Macrosomia-obesity-macrocephaly-ocular abnormalities syndrome	→399808	Male infertility due to NANOS1 mutation	3148	Malignant neurofibroma
141276	Macrostomia	399805	Male infertility with azoospermia or oligozoospermia due to single gene mutation	206538	Malignant non-dysgerminomatous germ cell tumor of ovary
83619	Macrostomia-preauricular tags-external ophthalmoplegia syndrome	→399805	Male infertility with normal virilization due to maturation arrest	99912	Malignant ovarian dysgerminoma
→182050	Macrothrombocytopenia with leukocyte inclusions	→399805	Male infertility with normal virilization due to meiosis defect	3286	Malignant paroxysmal ventricular tachycardia
220448	Macrothrombocytopenia with mitral valve insufficiency	→399805	Male infertility with teratozoospermia due to single gene mutation	252128	Malignant perineurioma
217335	MACS syndrome	399808	Male infertility with teratozoospermia due to single gene mutation	3148	Malignant peripheral nerve sheath tumor
137814	Macular amyloidosis	753	Male pseudohermaphroditism due to 5-alpha-reductase 2 deficiency	252212	Malignant peripheral nerve sheath tumor with rhabdomyosarcomatous differentiation
91494	Macular coloboma-cleft palate-hallux valgus syndrome	752	Male pseudohermaphroditism due to 17-beta-hydroxysteroid dehydrogenase 3 deficiency	213812	Malignant peripheral neuroectodermal tumor of the cervix uteri
98969	Macular corneal dystrophy				
79457	Maculopapular cutaneous mastocytosis	755	Male pseudohermaphroditism due to LH resistance or LHB deficiency	213630	Malignant peripheral neuroectodermal tumor of the corpus uteri
2457	MAD				
26791	MAD deficiency				
394532	MAD deficiency, mild type				
394529	MAD deficiency, severe neonatal type				
26791	MADD				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
168811	Malignant peritoneal mesothelioma	79113	Mandibulofacial dysostosis-microcephaly syndrome	2767	Maroteaux-Le Merrer-Bensahel syndrome
69077	Malignant rhabdoid tumor			950	Maroteaux-Malamut syndrome
3148	Malignant schwannoma	306682	Manganese intoxication	950	Maroteaux-Malamut syndrome
99916	Malignant Sertoli-Leydig cell tumor of ovary	306682	Manganese poisoning	1423	Maroteaux-Stanescu-Cousin syndrome
		306682	Manganism	1040	Maroteaux-Verloes-Stanescu syndrome
398987	Malignant teratoma of ovary	2717	Manitoba oculotrichoanal syndrome	1040	Maroteaux-Verloes-Stanescu syndrome
99868	Malignant thymoma	79327	Mannosyltransferase 1 deficiency	560	Marshall syndrome
252212	Malignant triton tumor	79326	Mannosyltransferase 2 deficiency	42642	Marshall syndrome with periodic fever
180242	Malignant tubal tumor	79321	Mannosyltransferase 6 deficiency	561	Marshall-Smith syndrome
180242	Malignant tumor of fallopian tubes	79328	Mannosyltransferase 7-9 deficiency	908	Martin-Bell syndrome
943	Malonic aciduria	79324	Mannosyltransferase 8 deficiency	85321	Martin-Probst syndrome
943	Malonyl-CoA decarboxylase deficiency	2459	Mansonelliasis	→293864	Martinez-Frías syndrome
		2459	Mansonellosis	466718	Martinique crinkled retinal pigment epitheliopathy
2229	Malouf syndrome	52416	Mantle cell lymphoma	1387	Martsof syndrome
99090	Malposition of the coronary ostium	52416	Mantle zone lymphoma	2466	MASA syndrome
→293843	Malpuech facial clefting syndrome	511	Maple syrup urine disease	→284963	MASS syndrome
→293843	Malpuech syndrome	→2712	Marashi-Gorlin syndrome	66661	Mast cell sarcoma
293843	Malpuech-Michels-Mingarelli-Carnevale syndrome	2785	Marble brain disease	101001	Mast syndrome
		228157	Marburg acute multiple sclerosis	2135	Mastocytosis-short stature-hearing loss syndrome
52417	MALT lymphoma	99826	Marburg hemorrhagic fever	3282	MAT
103907	Maltase-glucoamylase deficiency	99826	Marburg virus disease	168598	MAT deficiency
52417	MALToMa	221074	Marchiafava-Bignami disease	168598	MAT I/III deficiency
180275	Mammary Paget disease	447	Marchiafava-Micheli disease	254534	Maternal 14q32.2 hypermethylation syndrome
50920	Mammary polyadenomatosis	91412	Marcus-Gunn phenomenon	254528	Maternal 14q32.2 microdeletion syndrome
238744	Mammary-digital-nail syndrome	91412	Marcus-Gunn syndrome	275944	Maternal anti-Kell alloimmunization
397941	MAN1B1-CDG	2461	Marden-Walker syndrome	254528	Maternal del(14)(q32.2)
244310	Man5GlcNAc2-PP-Dol flippase deficiency	2460	Marden-Walker-like syndrome	2209	Maternal hyperphenylalaninemia
		1120	Mardini-Nyhan syndrome	2216	Maternal hyperthermia induced birth defects
141174	Mandibular arteriovenous malformation	558	Marfan syndrome	254528	Maternal monosomy 14q32.2
363649	Mandibular hypoplasia-deafness-progeroid syndrome	284963	Marfan syndrome type 1	2209	Maternal phenylketonuria
		284973	Marfan syndrome type 2	2209	Maternal PKU
246	Mandibulofacial dysostosis with postaxial limb anomalies	2462	Marfanoid craniosynostosis syndrome	411712	Maternal riboflavin deficiency
91412	Mandibulo-palpebral synkinesis-ptosis syndrome	2463	Marfanoid habitus-autosomal recessive intellectual disability syndrome	251009	Maternal uniparental disomy of chromosome 1
2457	Mandibuloacral dysplasia	→60030	Marfanoid habitus-craniosynostosis syndrome	96179	Maternal uniparental disomy of chromosome 2
90153	Mandibuloacral dysplasia with type A lipodystrophy	314041	Marfanoid habitus-inguinal hernia-advanced bone age syndrome	96180	Maternal uniparental disomy of chromosome 4
90154	Mandibuloacral dysplasia with type B lipodystrophy	2464	Marfanoid syndrome, De Silva type	96181	Maternal uniparental disomy of chromosome 6
443995	Mandibulofacial dysostosis with alopecia	→3253	Margarita island ectodermal dysplasia	96183	Maternal uniparental disomy of chromosome 9
245	Mandibulofacial dysostosis with preaxial limb anomalies	444	Marie Unna congenital hypotrichosis	97678	Maternal uniparental disomy of chromosome 13
861	Mandibulofacial dysostosis without limb anomalies	444	Marie Unna hereditary hypotrichosis	96184	Maternal uniparental disomy of chromosome 14
79113	Mandibulofacial dysostosis, Guion-Almeida type	101104	Marin-Amat syndrome	96185	Maternal uniparental disomy of chromosome 16
1131	Mandibulofacial dysostosis, Toriello type	559	Marinesco-Sjögren syndrome		
		2717	Marles syndrome		
357158	Mandibulofacial dysostosis-macroblepharon-macrostomia syndrome	2717	Marles-Greenberg-Persaud syndrome		
		583	Maroteaux-Lamy disease		

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96186	Maternal uniparental disomy of chromosome 20	93555	MC type III	210272	MdD
		254519	MCA due to 14q32.2 maternally expressed gene defect	210272	MdDS
96187	Maternal uniparental disomy of chromosome 21	1836		1836	MDK
		42	MCAD deficiency	238744	MDN syndrome
96188	Maternal uniparental disomy of chromosome 22	42	MCADD	363649	MDP syndrome
		300496	MCAHS type 2	3097	Meacham syndrome
261519	Maternal uniparental disomy of chromosome X	→56304	McAlister-Crane syndrome	3097	Meacham-Winn-Culler syndrome
		60040	MCAP	435438	MEAK
96186	Maternal UPD(20)	368	McArdle disease	370997	MEB disease with bilateral multicystic leucodystrophy
1349	Maternally-inherited cardiomyopathy and deafness	79140	MCC		
1349	Maternally-inherited cardiomyopathy and hearing loss	6	MCC deficiency	588	MEB syndrome
		85195	McCabe disease	98954	MECD
663	Maternally-inherited chronic progressive external ophthalmoplegia	6	MCCD	564	Meckel syndrome
		562	McCune-Albright syndrome	3032	Meckel syndrome type 7
663	Maternally-inherited CPEO	93686	MCD	564	Meckel-Gruber syndrome
		98969	MCD	3032	Meckel-like syndrome type 1
225	Maternally-inherited diabetes and deafness	1851	MC DK	70588	Meconium aspiration syndrome
		2471	McDonough syndrome	314376	Meconium ileus due to guanylate cyclase 2C deficiency
255210	Maternally-inherited infantile subacute necrotizing encephalopathy	→357225	McDowall syndrome		
		75327	MCDR1	93308	MED1
255210	Maternally-inherited Leigh disease	319640	MCDR2	93307	MED4
		36412	McDuffie hypocomplementemic urticarial vasculitis	93311	MED5
255210	Maternally-inherited Leigh syndrome	36412	McDuffie syndrome	98838	Med-DLBCL
		308425	MCEE deficiency	3453	MEDAC syndrome
254851	Maternally-inherited mitochondrial dystonia	158668	McGrath syndrome	2476	Medeira-Dennis-Donnai syndrome
		2473	McKusick-Kaufman syndrome	57196	Medial condensing osteitis of the clavicle
663	Maternally-inherited progressive external ophthalmoplegia	52416	MCL		
		320360	Maternally-inherited spastic paraplegia	59306	McLeod neuroacanthocytosis syndrome
320360	Maternally-inherited SPG			60040	MCM
2015	Mathieu-De Broca-Bony syndrome	60040	MCMTC	141239	Median cleft of the upper lip and maxilla
2470	Matthew-Wood syndrome	77298	MCOPS3	1993	Median cleft of the upper lip-corporum callosum lipoma-cutaneous polyps syndrome
552	Maturity-onset diabetes of the young	85275	MCOPS4		
		178364	MCOPS5	2699	Median nodule of the upper lip
293603	Maumenee corneal dystrophy	139471	MCOPS6	98838	Mediastinal diffuse large-cell lymphoma with sclerosis
141171	Maxillary arteriovenous malformation	2556	MCOPS7	63999	Mediastinal fibrosis
		3434	MCOPS8	370127	Medich giant platelet syndrome
1248	Maxillonasal dysostosis	2470	MCOPS9	370127	Medich macrothrombocytopenia
1248	Maxillonasal dysplasia	77299	MCOPS10	231	Medina worm disease
→182050	May-Hegglin anomaly	2512	MCPH	231	Medinensis
→182050	May-Hegglin syndrome	2001	McPherson-Clemens syndrome	231214	Mediterranean anemia
→182050	May-Hegglin thrombocytopenia	2999	McPherson-Hall syndrome	100025	Mediterranean lymphoma
3109	Mayer-Rokitansky-Küster-Hauser syndrome	319287	MCRCC	→2882	Mediterranean macrothrombocytopenia
		466718	MCRPE		
247775	Mayer-Rokitansky-Küster-Hauser syndrome type 1	59	MCT8 deficiency	83313	Mediterranean spotted fever
		809	MCTD		
2578	Mayer-Rokitansky-Küster-Hauser syndrome type 2	523	MCUL	42	Medium chain acyl-CoA dehydrogenase deficiency
		565	MD		
57782	Mazabraud syndrome	273	MD1	42	Medium chain acyl-coenzyme A dehydrogenase deficiency
251858	MBEN	258	MDC1A		
91138	MC	98893	MDC1B	171851	MEDNIK syndrome
71529	MC4R deficiency	→370953	MDC1C	3050	Medrano-Roldan syndrome
93554	MC type II	→370953	MDC1D		

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29073	Medullary plasmacytoma	90186	Meige disease	319547	Mendelian susceptibility to mycobacterial diseases due to complete interferon gamma receptor 2 deficiency
1309	Medullary sponge kidney	93964	Meige dystonia		
1332	Medullary thyroid carcinoma	90186	Meige lymphedema		
616	Medulloblastoma	93964	Meige syndrome	319558	Mendelian susceptibility to mycobacterial diseases due to complete interleukin 12B deficiency
251858	Medulloblastoma with extensive nodularity	→90186	Meige-like disease		
		314451	Meigs syndrome	319563	Mendelian susceptibility to mycobacterial diseases due to complete ISG15 deficiency
251883	Medulloepithelioma of the central nervous system	98868	Melanesian elliptocytosis		
		98868	Melanesian ovalocytosis		
98954	Meesmann corneal dystrophy	252206	Melanoma and neural system tumor syndrome	319600	Mendelian susceptibility to mycobacterial diseases due to partial interferon regulatory factor 8 deficiency
97252	Mega-cisterna magna				
66629	Megacolon-microcephaly syndrome	97338	Melanoma of soft tissue		
280671	Megaconial congenital muscular dystrophy	252206	Melanoma-astrocytoma syndrome	319600	Mendelian susceptibility to mycobacterial diseases due to partial IRF8 deficiency
238637	Megacystis-megaureter syndrome	404560	Melanoma-pancreatic cancer syndrome		
2241	Megacystis-microcolon-intestinal hypoperistalsis syndrome	79146	Melanosis diffusa congenita	319600	Mendelian susceptibility to mycobacterial diseases due to partial IRF8 deficiency
		79146	Melanosis universalis hereditaria		
2241	Megacystis-microcolon-intestinal hypoperistalsis-hydronephrosis syndrome	550	MELAS	319595	Mendelian susceptibility to mycobacterial diseases due to partial signal transducer and activator of transcription 1 deficiency
		87503	Meleda disease		
2604	Megaduodenum and/or megacystis	2482	Melhem-Fahl syndrome		
402023	Megakaryoblastic acute myeloid leukemia with t(1;22)(p13;q13)	31202	Melioidosis		
402023	Megakaryoblastic AML with t(1;22)(p13;q13)	2483	Melkersson-Rosenthal syndrome		
2478	Megalencephalic leukodystrophy	2484	Melnick-Needles osteodysplasty	319595	Mendelian susceptibility to mycobacterial diseases due to partial STAT1 deficiency
		2484	Melnick-Needles syndrome		
		2485	Melorheostosis	2494	Ménétrier disease
		1879	Melorheostosis with osteopoikilosis		
2478	Megalencephalic leukoencephalopathy with subcortical cysts	93571	Membranoproliferative glomerulonephritis type 2	3216	Mengel-Konigsmark syndrome
2477	Megalencephaly	652	MEN1	252046	Meningeal melanocytoma
60040	Megalencephaly-capillary malformation syndrome	653	MEN2	2495	Meningioma
		247698	MEN2A	→823	Meningocele
60040	Megalencephaly-capillary malformation-polymicrogyria syndrome	247709	MEN2B	33475	Meningococcal meningitis
		276152	MEN4	565	Menkes disease
		401973	MEND syndrome	565	Menkes syndrome
60040	Megalencephaly-cutis marmorata telangiectatica congenita syndrome	319552	Mendelian susceptibility to interleukin 12 receptor beta 1 deficiency	75858	Mental retardation-truncal obesity-retinal dystrophy-micropenis syndrome
2478	Megalencephaly-cystic leukodystrophy syndrome	99898	Mendelian susceptibility to mycobacterial diseases due to complete IFNgammaR1 deficiency	330021	Mercurialism
				330021	Mendelian susceptibility to mycobacterial diseases due to complete IFNgammaR2 deficiency
83473	Megalencephaly-polymicrogyria-postaxial polydactyly-hydrocephalus syndrome	319547	Mendelian susceptibility to mycobacterial diseases due to complete IL12B deficiency	330021	Mercury poisoning
457359	Megalencephaly-severe kyphoscoliosis-overgrowth syndrome	319558	Mendelian susceptibility to mycobacterial diseases due to complete IL12RB1 deficiency	79140	Merkel cell carcinoma
2479	Megalocornea-intellectual disability syndrome	319552	Mendelian susceptibility to mycobacterial diseases due to complete IL12RB1 deficiency	258	Merosin-negative congenital muscular dystrophy
238763	Megalocornea-spherophakia-secondary glaucoma syndrome			551	MERRF
50815	Mégarbané-Loiselet syndrome	99898	Mendelian susceptibility to mycobacterial diseases due to complete interferon gamma receptor 1 deficiency	54370	Mesangiocapillary glomerulonephritis
238637	Megaureter-megacystis syndrome			386	Mesenchymal hamartoma of liver
352328	MEGDEL syndrome			238593	Mesenteric lipogranuloma
3038	Mehes syndrome			238593	Mesenteric panniculitis
85282	MEHMO syndrome			99701	Mesial temporal lobe epilepsy with hippocampal sclerosis
2196	Meier-Blumberg-Imahorn syndrome			295004	Mesoaxial polydactyly of fingers
2554	Meier-Gorlin syndrome			295173	Mesoaxial polydactyly of fingers, bilateral
				295171	Mesoaxial polydactyly of fingers, unilateral
				295010	Mesoaxial polydactyly of toes

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295185	Mesoaxial polydactyly of toes, bilateral	2502	Metaphyseal dysostosis-intellectual disability-conductive deafness syndrome	308425	Methylmalonic aciduria due to methylmalonyl-CoA epimerase deficiency
295183	Mesoaxial polydactyly of toes, unilateral			308425	Methylmalonic aciduria due to methylmalonyl-CoA racemase deficiency
157801	Mesoaxial synostotic syndactyly with phalangeal reduction	→175	Metaphyseal dysplasia without hypotrichosis	308425	Methylmalonic aciduria due to methylmalonyl-CoA racemase deficiency
95443	Mesocardia	85188	Metaphyseal dysplasia, Braun-Tinschert type	280183	Methylmalonic aciduria due to transcobalamin receptor defect
289	Mesodermic dysplasia	3005	Metaphyseal dysplasia, Pyle type	26	Methylmalonic aciduria with homocystinuria
2496	Mesomelia-synostoses syndrome	2504	Metaphyseal dysplasia-maxillary hypoplasia-brachydacty syndrome	79282	Methylmalonic aciduria with homocystinuria, type cbIC
2496	Mesomelia-synostoses syndrome, Verloes-David-Pfeiffer type	213531	Metaplastic carcinoma of the breast	79283	Methylmalonic aciduria with homocystinuria, type cbID
2632	Mesomelic dwarfism, Langer type	2635	Metatropic dwarfism	79284	Methylmalonic aciduria with homocystinuria, type cbIF
2633	Mesomelic dwarfism, Nievergelt type	2635	Metatropic dysplasia	369955	Methylmalonic aciduria with homocystinuria, type cbIJ
2634	Mesomelic dwarfism, Reinhardt-Pfeiffer type	88639	Methacrylic aciduria	369962	Methylmalonic aciduria with homocystinuria, type cbIX
		31825	Methanol poisoning	29	Mevalonic aciduria
2631	Mesomelic dwarfism-cleft palate-camptodactyly syndrome	1923	Methimazole embryofetopathy	2710	Meyer-Schwickerath syndrome
		1923	Methimazole/carbimazole embryofetopathy	443995	MFDA
97360	Mesomelic dwarfism-small genitalia syndrome	1923	Methimazole/carbimazole embryopathy	79113	MFDM syndrome
85170	Mesomelic dysplasia with absent fibulas and triangular tibias	168598	Methionine adenosyltransferase deficiency	558	MFS
				2710	Meyer-Schwickerath syndrome
2496	Mesomelic dysplasia with acral synostoses, Verloes-David-Pfeiffer type	86904	Methotrexate-associated lymphoproliferative disorders	284963	MFS1
				1917	Methyl mercury antenatal infection
1836	Mesomelic dysplasia, Kantaputra type	622	Methylcobalamin deficiency	67046	MGA1
85170	Mesomelic dysplasia, Savarirayan type	308380	Methylcobalamin deficiency type cbIDv1	111	MGA2
		2169	Methylcobalamin deficiency type cbIE	67047	MGA3
171690	Metabolic myopathy due to lactate transporter defect	2170	Methylcobalamin deficiency type cbIG	67048	MGA4
2499	Metachondromatosis	395	Methylene tetrahydrofolate reductase deficiency	66634	MGA5
512	Metachromatic leukodystrophy			445038	MGA7
309271	Metachromatic leukodystrophy, adult form	308425	Methylmalonic acidemia due to methylmalonyl-CoA epimerase deficiency	79329	MGAT2-CDG
				→182050	MHA
309263	Metachromatic leukodystrophy, juvenile form	308425	Methylmalonic acidemia due to methylmalonyl-CoA racemase deficiency	443162	MHAC
				391417	MHBD deficiency
309256	Metachromatic leukodystrophy, late infantile form	308425	Methylmalonic acidemia due to methylmalonyl-CoA racemase deficiency	391428	MHBD deficiency, classic type
				391428	MHBD deficiency, infantile type
1240	Metaphyseal acroscaphodysplasia	26	Methylmalonic acidemia with homocystinuria	391457	MHBD deficiency, neonatal type
1040	Metaphyseal anadysplasia			99826	MHF
33067	Metaphyseal chondrodysplasia, Jansen type	79284	Methylmalonic acidemia with homocystinuria type cbIF	386	MHL
166038	Metaphyseal chondrodysplasia, Kaitila type	79282	Methylmalonic acidemia with homocystinuria, type cbIC	79651	mHPA
				79283	Methylmalonic acidemia with homocystinuria, type cbID
175	Metaphyseal chondrodysplasia, McKusick type	79283	Methylmalonic acidemia with homocystinuria, type cbID	294016	MIC-CM syndrome
174	Metaphyseal chondrodysplasia, Schmid type	369955	Methylmalonic acidemia with homocystinuria, type cbIJ	→293843	Michels syndrome
				163937	MICPCH
2501	Metaphyseal chondrodysplasia, Spahr type	369962	Methylmalonic acidemia with homocystinuria, type cbIX	2510	Micro syndrome
				2511	Microbrachycephaly-ptosis-cleft lip syndrome
166035	Metaphyseal chondrodysplasia-retinitis pigmentosa syndrome	280183	Methylmalonic acidemia, TCb1R type	2512	Microcephalia vera
				280183	Methylmalonic acidemia, TCb1R type
99646	Metaphyseal chondromatosis with D-2-hydroxyglutaric aciduria	280183	Methylmalonic acidemia, TCb1R type	2637	Microcephalic osteodysplastic primordial dwarfism type II

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2636	Microcephalic osteodysplastic primordial dwarfism types I and III	294016	Microcephaly-cutaneous capillary malformation syndrome	1305	Microcephaly-oculo-digito-esophageal-duodenal syndrome
2636	Microcephalic osteodysplastic primordial dwarfism, Taybi-Linder type	2533	Microcephaly-deafness-intellectual disability syndrome	391641	Microcephaly-oculo-digito-esophageal-duodenal syndrome type 1
468631	Microcephalic primordial dwarfism due to RTTN deficiency	137653	Microcephaly-digital anomalies-intellectual disability syndrome	171703	Microcephaly-polymicrogyria-corpora callosa agenesis syndrome
329228	Microcephalic primordial dwarfism due to ZNF335 deficiency	1305	Microcephaly-digital anomalies-normal intelligence syndrome	2519	Microcephaly-seizures-intellectual disability-heart disease syndrome
319671	Microcephalic primordial dwarfism, Alazami type	391641	Microcephaly-digital anomalies-normal intelligence syndrome type 1	423306	Microcephaly-short stature-intellectual disability-facial dysmorphism syndrome
319675	Microcephalic primordial dwarfism, Dauber type	391646	Microcephaly-digital anomalies-normal intelligence syndrome type 2	397951	Microcephaly-thin corpus callosum-intellectual disability syndrome
2643	Microcephalic primordial dwarfism, Toriello type	217026	Microcephaly-facio-cardio-skeletal syndrome, Hadziselimovic type	2670	Microcoria-congenital nephrosis syndrome
329228	Microcephalic primordial dwarfism, Walsh type	217026	Microcephaly-faciocardioskeletal syndrome	2535	Microcornea-corectopia-macular hypoplasia syndrome
436182	Microcephalic primordial dwarfism-insulin resistance syndrome	2172	Microcephaly-glomerulonephritis-marfanoid habitus syndrome	2536	Microcornea-glaucoma-absent frontal sinuses syndrome
240760	Microcephaly and chromosomal instability without immunodeficiency	2065	Microcephaly-hiatus hernia-nephrotic syndrome	369970	Microcornea-myopic chorioretinal atrophy-telecanthus syndrome
2512	Microcephaly vera	2558	Microcephaly-hypergonadotropic hypogonadism-short stature syndrome	231736	Microcornea-posterior megalolenticonus-persistent fetal vasculature-coloiboma syndrome
2513	Microcephaly-albinism-digital anomalies syndrome	3132	Microcephaly-hypogammaglobulinemia-abnormal immunity syndrome	263347	Microcornea-rod-cone dystrophy-cataract-posterior staphyloma syndrome
3433	Microcephaly-brachydactyly-kyphoscoliosis syndrome	647	Microcephaly-immunodeficiency-lymphoreticuloma syndrome	98956	Microcystic corneal dystrophy
2523	Microcephaly-brain defect-spasticity-hypernatremia syndrome	137658	Microcephaly-intellectual disability-phalangeal and neurological anomalies syndrome	79490	Microcystic infiltrating lymphatic malformation
294016	Microcephaly-capillary malformation syndrome	457351	Microcephaly-intellectual disability-sensorineural deafness-epilepsy-abnormal muscle tone syndrome	79490	Microcystic lymphangioma
2516	Microcephaly-cardiac defect-lung malsegmentation syndrome	457351	Microcephaly-intellectual disability-sensorineural hearing loss-epilepsy-abnormal muscle tone syndrome	79490	Microcystic lymphatic malformation
2515	Microcephaly-cardiomyopathy syndrome	1305	Microcephaly-intellectual disability-tracheoesophageal fistula syndrome	83642	Microcytic anemia with liver iron overload
329332	Microcephaly-cerebellar hypoplasia-cardiac conduction defect syndrome	391641	Microcephaly-intellectual disability-tracheoesophageal fistula syndrome type 1	77301	Microdeletion 9q22.3
329332	Microcephaly-cerebellar hypoplasia-congenital heart conduction defect syndrome	391646	Microcephaly-intellectual disability-tracheoesophageal fistula syndrome type 2	567	Microdeletion 22q11.2
434179	Microcephaly-cerebral malformation-orofacioidigital syndrome	1229	Microcephaly-intracranial calcification-intellectual disability syndrome	90024	Microdontia-type I microtia-deafness syndrome
2522	Microcephaly-cervical spine fusion anomalies syndrome	2526	Microcephaly-lymphedema-chorioretinopathy syndrome	101081	Microduplication 17p12
2521	Microcephaly-cleft palate syndrome	2528	Microcephaly-microcornea syndrome, Seemanova type	217377	Microduplication Xp11.22-p11.23 syndrome
423894	Microcephaly-complex motor and sensory axonal neuropathy syndrome	3434	Microcephaly-microphthalmia-ectrodactyly of lower limbs-prognathism syndrome	280200	Microform holoprosencephaly
2508	Microcephaly-corpora callosa agenesis-abnormal genitalia syndrome			280200	Microform HPE
457284	Microcephaly-corpora callosa hypoplasia-intellectual disability-facial dysmorphism syndrome			2538	Microgastria-limb reduction defect syndrome
				1388	Micrognathia digital syndrome
				476126	Micrognathia-recurrent infections-behavioral abnormalities-mild intellectual disability syndrome
				1083	Microlissencephaly
				89844	Microlissencephaly type A
				50810	Microlissencephaly-micromelia syndrome
				2641	Micromelic dwarfism, Fryns type

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
93329	Micromelic dysplasia-dislocation of radius syndrome	93926	Middle interhemispheric fusion variant	98832	Minimally differentiated acute myeloblastic leukemia
139471	Microphthalmia with brain and digit anomalies	93926	Middle interhemispheric variant of holoprosencephaly	822	Minkowski-Chauffard disease
98938	Microphthalmia with colobomatous cyst	141288	Midline cervical cleft	1918	Minoxidil antenatal infection
1104	Microphthalmia with facial clefting	95443	Midline heart	94125	MIRAS
1106	Microphthalmia with limb anomalies	93926	Midline interhemispheric variant of holoprosencephaly	→193	Mirhosseini-Holmes-Walton syndrome
2556	Microphthalmia with linear skin defects syndrome	2557	Mietens syndrome	295010	Mirror foot
568	Microphthalmia, Lenz type	2867	Mievis-Verellen-Dumoulin syndrome	295185	Mirror foot, bilateral
85275	Microphthalmia-ankyloblepharon-intellectual disability syndrome	293181	Migrating partial epilepsy of infancy	295183	Mirror foot, unilateral
98938	Microphthalmia-anophthalmia-coloboma syndrome	293181	Migrating partial seizures of infancy	295004	Mirror hand
77299	Microphthalmia-brain atrophy syndrome	504	Migratory myiasis	295173	Mirror hand, bilateral
424099	Microphthalmia-coloboma-rhizomelic skeletal dysplasia	93926	MIH	295171	Mirror hand, unilateral
2556	Microphthalmia-dermal aplasia-sclerocornea syndrome	93926	MIH type HPE	2378	Mirror hands and feet-nasal defects syndrome
→2510	Microphthalmia-intellectual disability syndrome	93926	MIHF	3004	Mirror polydactyly-vertebral segmentation-limbs defects syndrome
2547	Microphthalmia-microtia-fetal akinesia syndrome	93926	MIHV	319308	MiT family translocation renal cell carcinoma
2705	Microphthalmia-optic nerve aplasia syndrome	2558	Mikati-Najjar-Sahli syndrome	293822	MITF-related melanoma and renal cell carcinoma predisposition syndrome
251279	Microphthalmia-retinitis pigmentosa-foveoschisis-optic disc drusen syndrome	79078	Mikulicz disease	134	Mitochondrial acetoacetyl-coenzyme A thiolase deficiency
727	Micropolyangiitis	314918	Mild Canavan disease	353217	Mitochondrial aspartate-glutamate carrier 1 deficiency
727	Microscopic polyangiitis	169799	Mild factor IX deficiency	225	Mitochondrial diabetes
727	Microscopic polyarteritis	169808	Mild factor VIII deficiency	352470	Mitochondrial DNA deletion syndrome with limb-girdle weakness
2551	Microspherophakia-metaphyseal dysplasia syndrome	169808	Mild hemophilia A	352470	Mitochondrial DNA deletion syndrome with progressive myopathy
2552	Microsporidiosis	169799	Mild hemophilia B	1933	Mitochondrial DNA depletion syndrome, encephalomyopathic form with methylmalonic aciduria
83463	Microtia	79651	Mild HPA	255235	Mitochondrial DNA depletion syndrome, encephalomyopathic form with renal tubulopathy
2306	Microtia-aortic arch syndrome	79651	Mild hyperphenylalaninemia	369897	Mitochondrial DNA depletion syndrome, encephalomyopathic form with variable craniofacial anomalies
139450	Microtia-eye coloboma-imperforation of the nasolacrimal duct syndrome	171439	Mild nemaline myopathy	279934	Mitochondrial DNA depletion syndrome, hepatocerebral form due to DGUOK deficiency
289522	Microtriplication 11q24.1	216796	Mild osteogenesis imperfecta	363534	Mitochondrial DNA depletion syndrome, hepatocerebrorenal form
2290	Microvillous inclusion disease	247815	Mild peroxisomal disorder due to PEX10 deficiency	254875	Mitochondrial DNA depletion syndrome, myopathic form
2290	Microvillus inclusion disease	79253	Mild phenylketonuria	352447	Mitochondrial DNA maintenance syndrome due to MGME1 deficiency
166430	Micturation-induced seizures	411536	Mild phosphoribosylpyrophosphate synthetase superactivity	1194	Mitochondrial encephalo-cardio-myopathy due to F1Fo ATPase deficiency
1456	Mid-aortic dysplastic syndrome	79253	Mild PKU		
1456	Mid-aortic syndrome	411536	Mild PRPP synthetase superactivity		
228299	Mid-dermal elastolysis	411536	Mild PRPS1 superactivity		
1456	Midaortic syndrome	93279	Mild spondyloepiphyseal dysplasia due to COL2A1 mutation with early-onset osteoarthritis		
2556	MIDAS syndrome	246	Miller syndrome		
225	MIDD	531	Miller-Dieker syndrome		
1456	Middle aortic syndrome	98919	Miller-Fisher syndrome		
100084	Middle ear neuroendocrine tumor	94091	Mills syndrome		
		79452	Milroy disease		
		→79452	Milroy-like disease		
		255210	MILS		
		1917	Minamata disease		
		457485	MINDS syndrome		
		757	Mineralocorticoid resistant hyperkalemia		
		→293843	Mingarelli syndrome		
		352734	Minimal pigment oculocutaneous albinism type 1		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
1194	Mitochondrial encephalo-cardio-myopathy due to isolated ATP synthase deficiency	298	Mitochondrial neurogastrointestinal encephalomyopathy	399096	Miyoshi muscular dystrophy type 3
				45448	Miyoshi myopathy
1194	Mitochondrial encephalo-cardio-myopathy due to isolated mitochondrial respiratory chain complex V deficiency	90641	Mitochondrial non-syndromic neurosensory deafness	98757	MJD
				565	MK
238329	Mitochondrial encephalomyopathy due to combined oxidative phosphorylation defect 6	168609	Mitochondrial non-syndromic neurosensory deafness with susceptibility to aminoglycoside exposure	423461	ML 3 alpha/beta
				423470	ML 3 gamma
238329	Mitochondrial encephalomyopathy due to COXPD6	168609	Mitochondrial non-syndromic neurosensory hearing loss with susceptibility to aminoglycoside exposure	423461	ML III alpha/beta
				423470	ML III gamma
550	Mitochondrial encephalomyopathy, lactic acidosis and stroke-like episodes	90641	Mitochondrial non-syndromic sensorineural deafness	2598	MLASA
				2478	MLC
1933	Mitochondrial encephalomyopathy-aminoacidopathy syndrome	168609	Mitochondrial non-syndromic sensorineural deafness with susceptibility to aminoglycoside exposure	2526	MLCRD
				512	MLD
280288	Mitochondrial HSP60 chaperonopathy	168609	Mitochondrial non-syndromic sensorineural deafness with susceptibility to aminoglycoside exposure	309271	MLD, adult form
				309263	MLD, juvenile form
314637	Mitochondrial hypertrophic cardiomyopathy with lactic acidosis due to MTO1 deficiency	168609	Mitochondrial non-syndromic sensorineural hearing loss with susceptibility to aminoglycoside exposure	309256	MLD, late infantile form
				59306	MLS
168609	Mitochondrial isolated neurosensory deafness with susceptibility to aminoglycoside exposure	447784	Mitochondrial pyruvate carrier deficiency	2556	MLS syndrome
		254881	Mitochondrial spinocerebellar ataxia with epilepsy	464321	MLT
168609	Mitochondrial isolated neurosensory hearing loss with susceptibility to aminoglycoside exposure	746	Mitochondrial trifunctional protein deficiency	369970	MMCAT syndrome
		1205	Mitral atresia	598	MmD
168609	Mitochondrial isolated sensorineural deafness with susceptibility to aminoglycoside exposure	3238	Mitral regurgitation-deafness-skeletal anomalies syndrome	399096	MMD3
		99062	Mitral valve agenesis	3434	MMEP syndrome
168609	Mitochondrial isolated sensorineural hearing loss with susceptibility to aminoglycoside exposure	→284963	Mitral valve-aorta-skeleton-skin syndrome	592	MMF
		295012	Mitten hand	268249	MMF embryopathy
289560	Mitochondrial membrane protein-associated neurodegeneration	90036	Mixed AIHA	2241	MMIHS
		809	Mixed connective tissue disease	213512	MMMT of the ovary
2598	Mitochondrial myopathy and sideroblastic anemia	91138	Mixed cryoglobulinemia	641	MMN
		93555	Mixed cryoglobulinemia type III	641	MMNCB
254864	Mitochondrial myopathy with reversible complex IV deficiency	458792	Mixed cystic lymphangioma	137867	MMND
		458792	Mixed cystic lymphatic malformation	293181	MMPEI
254864	Mitochondrial myopathy with reversible COX deficiency	180234	Mixed germ cell tumor	293181	MMPSI
		252021	Mixed germ cell tumor of central nervous system	2479	MMR syndrome
254864	Mitochondrial myopathy with reversible cytochrome C oxidase deficiency	252021	Mixed germ cell tumor of CNS	1305	MMT
		213610	Mixed Müllerian cancer of corpus uteri	391641	MMT type 1
550	Mitochondrial myopathy, encephalopathy, lactic acidosis and stroke-like episodes	251656	Mixed oligoastrocytoma	391646	MMT type 2
		2785	Mixed renal tubular acidosis	298	MNGIE
2597	Mitochondrial myopathy-lactic acidosis-deafness syndrome	2785	Mixed RTA	565	MNK
		1879	Mixed sclerosing bone dystrophy	251656	MOA
2597	Mitochondrial myopathy-lactic acidosis-hearing loss syndrome	324364	Mixed sclerosing bone dystrophy with extra-skeletal manifestations	77299	MOBA syndrome
		90036	Mixed-type autoimmune hemolytic anemia	570	Möbius syndrome
				99732	MOCOD
				308386	MOCOD type A
				308393	MOCOD type B
				308400	MOCOD type C
				1305	MODED syndrome
				391641	MODED syndrome type 1
				90056	Moderate and severe traumatic brain injury
				178145	Moderate multimicore disease with hand involvement
				169796	Moderately severe factor IX deficiency
				169805	Moderately severe factor VIII deficiency

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
169805	Moderately severe hemophilia A	261349	Monosomy 2p15p16.1	96149	Monosomy 12qter
169796	Moderately severe hemophilia B	163693	Monosomy 2p21	412035	Monosomy 13q12.3
263335	Moderately-differentiated thymic neuroendocrine carcinoma	228402	Monosomy 2q23.1	1587	Monosomy 13q14
		1617	Monosomy 2q24	1590	Monosomy 13q32
552	MODY	251014	Monosomy 2q31.1	96168	Monosomy 13q34
93111	MODY5	251019	Monosomy 2q32	261120	Monosomy 14q11.2
570	Moebius syndrome	251019	Monosomy 2q32-q33	261144	Monosomy 14q12
2560	Moebius syndrome-axonal neuropathy-hypogonadotropic hypogonadism syndrome	251019	Monosomy 2q32q33	→3157	Monosomy 14q22
		251028	Monosomy 2q33.1	264200	Monosomy 14q22-q23
		1001	Monosomy 2q37-qter	264200	Monosomy 14q22q23
1420	Moerman-Vandenbergh-Fryns syndrome	435638	Monosomy 3p25.3	401935	Monosomy 14q24.1q24.3
		1620	Monosomy 3pter	261183	Monosomy 15q11.2
3198	Moersch-Woltman syndrome	1621	Monosomy 3q13	199318	Monosomy 15q13.3
2549	Moeschler-Clarren syndrome	356947	Monosomy 3q26-q27	261190	Monosomy 15q14
79330	MOGS-CDG	356947	Monosomy 3q26q27	94065	Monosomy 15q24
2751	Mohr syndrome	65286	Monosomy 3q29	1596	Monosomy 15q26
2753	Mohr-Majewski syndrome	65286	Monosomy 3qter	261211	Monosomy 16p11.2-p12.2
52368	Mohr-Tranebjaerg syndrome	238750	Monosomy 4q21	261211	Monosomy 16p11.2p12.2
99927	Molar pregnancy	96145	Monosomy 4qter	261236	Monosomy 16p13.11
1433	Moloney syndrome	281	Monosomy 5p	352629	Monosomy 16q24.1
397973	MOMES syndrome	228384	Monosomy 5q14.3	261250	Monosomy 16q24.3
2563	MOMO syndrome	314655	Monosomy 5q31.3	531	Monosomy 17p13.3
371428	MONA spectrum	1627	Monosomy 5q35	97685	Monosomy 17q11
573	Monilethrix	251046	Monosomy 6p22	261265	Monosomy 17q12
573	Moniliform hair syndrome	96125	Monosomy 6p25	363958	Monosomy 17q21.31
319254	Monkey disease	171829	Monosomy 6q16	261279	Monosomy 17q23.1-q23.2
319254	Monkey fever	251056	Monosomy 6q25	261279	Monosomy 17q23.1q23.2
3057	Monoamine oxidase A deficiency	96126	Monosomy 7pter	1597	Monosomy 17qter
59	Monocarboxylate transporter 8 deficiency	904	Monosomy 7q11.23	1598	Monosomy 18p
		251061	Monosomy 7q31	1600	Monosomy 18q
228423	Monocyte-B-natural killer-dendritic cell deficiency syndrome	1636	Monosomy 7qter	254346	Monosomy 19p13.12
		251066	Monosomy 8p11.2	357001	Monosomy 19p13.13
228423	Monocytopenia and mycobacterial infection syndrome	251071	Monosomy 8p23.1	217346	Monosomy 19q13.11
		2496	Monosomy 8q13	261295	Monosomy 20p12.3
228423	Monocytopenia with susceptibility to infections	284160	Monosomy 8q21.11	313781	Monosomy 20p13
		178303	Monosomy 8q22.1	444051	Monosomy 20q11
99885	Monogenic diabetes of infancy	502	Monosomy 8q24.1	261311	Monosomy 20q13.33
228423	MonoMAC	261112	Monosomy 9p	96152	Monosomy 20qter
65684	Monomelic amyotrophy	324313	Monosomy 9p13	574	Monosomy 21
86870	Monomorphic NK-cell lymphoma	1642	Monosomy 9pter	261323	Monosomy 21q22.11-q22.12
2565	Mononen-Karnes-Senac syndrome	77301	Monosomy 9q22.3	261323	Monosomy 21q22.11q22.12
		401923	Monosomy 9q31.1q31.3	268261	Monosomy 21q22.13-q22.2
2901	Mononeuritis multiplex with brachial predilection	284169	Monosomy 10p11.21p12.31	268261	Monosomy 21q22.13q22.2
293948	Monosomy 1p21.3	1580	Monosomy 10pter	96123	Monosomy 22
401986	Monosomy 1p31p32	276413	Monosomy 10q22.3q23.3	567	Monosomy 22q11
456298	Monosomy 1p35.2	96148	Monosomy 10qter	48652	Monosomy 22q13
1606	Monosomy 1p36	893	Monosomy 11p13	99226	Monosomy X
1606	Monosomy 1pter	444002	Monosomy 11q22.2-q22.3	261476	Monosomy Xp21
250989	Monosomy 1q21.1	444002	Monosomy 11q22.2q22.3	93277	Monostotic fibrous dysplasia
250999	Monosomy 1q41-q42	2308	Monosomy 11qter	158003	Montgomery syndrome
250999	Monosomy 1q41q42	313884	Monosomy 12p12.1	→969	Moore-Federman syndrome
238769	Monosomy 1q44	94063	Monosomy 12q14	2637	MOPD type II
36367	Monosomy 1qter	289513	Monosomy 12q15q21.1	2636	MOPD types I and III
261349	Monosomy 2p15-p16.1				

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52056	Morava-Mehes syndrome	280679	Moyamoya angiopathy-short stature-facial dysmorphism-hypergonadotropic hypogonadism syndrome	217093	MPSIIB
77296	Morgagni-Stewart-Morel syndrome			581	MPSIII
75858	MORM syndrome			79269	MPSIIIA
35737	Morning glory syndrome			79270	MPSIIIB
582	Morquio disease	2573	Moyamoya disease	79271	MPSIIIC
309297	Morquio disease type A	401945	Moyamoya disease with early-onset achalasia	79272	MPSIIID
309310	Morquio disease type B			93474	MPSIS
2570	Morse-Rawnsley-Sargent syndrome	280679	Moyamoya disease-short stature-facial dysmorphism-hypergonadotropic hypogonadism	582	MPSIV
83467	Morvan syndrome			309297	MPSIVA
83467	Morvan's fibrillary chorea			309310	MPSIVB
329813	Mosaic genome-wide paternal uniparental disomy			2574	Moynahan syndrome
329813	Mosaic genome-wide paternal UPD	352734	MP OCA type 1	583	MPSVI
99228	Mosaic monosomy X	727	MPA	276212	MPSVI, rapidly progressing
96193	Mosaic paternal uniparental disomy of chromosome 11	289560	MPAN	276223	MPSVI, slowly progressing
1692	Mosaic trisomy 1	59135	MPD1	584	MPSVII
1723	Mosaic trisomy 2	399086	MPD3	99967	MRCLS
100071	Mosaic trisomy 3	79323	MPDU1-CDG	263347	MRCS syndrome
96059	Mosaic trisomy 4	293181	MPEI	67045	MRGH
96060	Mosaic trisomy 5	54370	MPGN	3109	MRKH syndrome
1747	Mosaic trisomy 7	79319	MPI-CDG	247775	MRKH syndrome type 1
96061	Mosaic trisomy 8	79253	mPKU	2578	MRKH syndrome type 2
99776	Mosaic trisomy 9	3148	MPNST	→457240	MRX35
96063	Mosaic trisomy 10	252212	MPNST with rhabdomyosarcomatous differentiation	85274	MRXS7
1698	Mosaic trisomy 12			85324	MRXS9
1703	Mosaic trisomy 14	2587	MPO deficiency	93952	MRXSH
1706	Mosaic trisomy 15	231736	MPPC syndrome	2598	MSA
1708	Mosaic trisomy 16	83473	MPPH syndrome	102	MSA
1711	Mosaic trisomy 17	579	MPS1	227510	MSA, cerebellar type
1724	Mosaic trisomy 20	93473	MPS1H	98933	MSA, parkinsonian type
96068	Mosaic trisomy 22	93476	MPS1H/S	227510	MSA-c
1052	Mosaic variegated aneuploidy syndrome	93474	MPS1S	98933	MSA-p
54057	Moschcowitz disease	580	MPS2	1879	MSBD syndrome
2717	MOTA syndrome	217085	MPS2A	254881	MSCAE
254516	Motor developmental delay due to 14q32.2 paternally expressed gene defect	217093	MPS2B	585	MSD
3347	Mounier-Kühn syndrome	581	MPS3	2619	Mseleni joint disease
83595	Mountain fever	79269	MPS3A	480536	MSH3-related AFAP
83595	Mountain tick fever	79270	MPS3B	480536	MSH3-related attenuated familial adenomatous polyposis
2572	Mousa-Al Din-Al Nassar syndrome	79271	MPS3C	480536	MSH3-related attenuated familial polyposis coli
324972	Mouth and genital ulcers with inflamed cartilage	79272	MPS3D	480536	MSH3-related attenuated FAP
2152	Mowat-Wilson syndrome	582	MPS4	1309	MSK
261537	Mowat-Wilson syndrome due to 2q22 microdeletion	309297	MPS4A	99898	MSMD due to complete IFNgamma1 deficiency
261552	Mowat-Wilson syndrome due to a ZEB2 point mutation	309310	MPS4B	319547	MSMD due to complete IFNgamma2 deficiency
261537	Mowat-Wilson syndrome due to del(2)q(22)	583	MPS6	319558	MSMD due to complete IL12B deficiency
261537	Mowat-Wilson syndrome due to monosomy 2q22	276212	MPS6, rapidly progressing	319552	MSMD due to complete IL12RB1 deficiency
		276223	MPS6, slowly progressing	99898	MSMD due to complete interferon gamma receptor 1 deficiency
		584	MPS7	319547	MSMD due to complete interferon gamma receptor 2 deficiency
		67041	MPS9		
		579	MPSI		
		293181	MPSI		
		93473	MPSIH		
		93476	MPSIH/S		
		580	MPSII		
		217085	MPSIIA		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
319552	MSMD due to complete interleukin 12 receptor beta 1 deficiency	577	Mucopolipidosis type III	276223	Mucopolysaccharidosis type VI, slowly progressing
		423461	Mucopolipidosis type III alpha/beta		
319558	MSMD due to complete interleukin 12B deficiency	423470	Mucopolipidosis type III gamma	584	Mucopolysaccharidosis type VII
		578	Mucopolipidosis type IV	73263	Mucormycosis
319563	MSMD due to complete ISG15 deficiency	579	Mucopolysaccharidosis type 1	52417	Mucosa-associated lymphatic tissue lymphoma
		93473	Mucopolysaccharidosis type 1H		
319600	MSMD due to partial interferon regulatory factor 8 deficiency	93476	Mucopolysaccharidosis type 1H/S	52417	Mucosa-associated lymphoid tissue lymphoma
		93474	Mucopolysaccharidosis type 1S		
319600	MSMD due to partial IRF8 deficiency	580	Mucopolysaccharidosis type 2	46486	Mucosal pemphigoid
				585	Mucosulfatidosis
319595	MSMD due to partial signal transducer and activator of transcription 1 deficiency	217093	Mucopolysaccharidosis type 2, attenuated form	46486	Mucosynechial pemphigoid
				46486	Mucous membrane pemphigoid
319595	MSMD due to partial STAT1 deficiency	217085	Mucopolysaccharidosis type 2, severe form	586	Mucoviscidosis
157801	MSSD	217085	Mucopolysaccharidosis type 2A	53271	Muenke syndrome
65748	MSSE	217093	Mucopolysaccharidosis type 2B	444	MUHH
511	MSUD	581	Mucopolysaccharidosis type 3	587	Muir-Torre syndrome
1332	MTC	79269	Mucopolysaccharidosis type 3A	2576	MULIBREY dwarfism
		79270	Mucopolysaccharidosis type 3B	2576	MULIBREY nanism
352470	mtDNA deletion syndrome with limb-girdle weakness	79271	Mucopolysaccharidosis type 3C		
		79272	Mucopolysaccharidosis type 3D	247768	Müllerian aplasia and hyperandrogenism
352470	mtDNA deletion syndrome with progressive myopathy	582	Mucopolysaccharidosis type 4		
		309297	Mucopolysaccharidosis type 4A	1655	Müllerian derivatives-lymphangiectasia-polydactyly syndrome
1933	mtDNA depletion syndrome, encephalomyopathic form with methylmalonic aciduria	309310	Mucopolysaccharidosis type 4B		
		583	Mucopolysaccharidosis type 6	2491	Müllerian duct anomalies-limb anomalies syndrome
255235	mtDNA depletion syndrome, encephalomyopathic form with renal tubulopathy	276212	Mucopolysaccharidosis type 6, rapidly progressing		
		276223	Mucopolysaccharidosis type 6, slowly progressing	2578	Müllerian duct aplasia-renal dysplasia-cervical somite anomalies syndrome
369897	mtDNA depletion syndrome, encephalomyopathic form with variable craniofacial anomalies	584	Mucopolysaccharidosis type 7	247768	Müllerian duct failure and hyperandrogenism
		67041	Mucopolysaccharidosis type 9		
363534	mtDNA depletion syndrome, hepatocerebrorenal form	579	Mucopolysaccharidosis type I	2774	Multicentric carpo-tarsal osteolysis with or without nephropathy
		93473	Mucopolysaccharidosis type IH	93686	Multicentric Castleman disease
254875	mtDNA depletion syndrome, myopathic form	93476	Mucopolysaccharidosis type IH/S	93686	Multicentric giant lymph node hyperplasia
		580	Mucopolysaccharidosis type II		
352447	mtDNA maintenance syndrome due to MGME1 deficiency	217093	Mucopolysaccharidosis type II, attenuated form	371428	Multicentric osteolysis-nodulosis-arthropathy spectrum
395	MTHFR deficiency				
252212	MTT	217085	Mucopolysaccharidosis type II, severe form	85196	Multicentric osteolysis-nodulosis-arthropathy syndrome
100024	mu-HCD	217085	Mucopolysaccharidosis type IIA	139436	Multicentric reticulohistiocytosis
100024	Mu-heavy chain disease	217093	Mucopolysaccharidosis type IIB	1851	Multicystic dysplastic kidney
398961	Mucinous adenocarcinoma of ovary	581	Mucopolysaccharidosis type III	168816	Multicystic mesothelioma
391723	Mucinous adenocarcinoma of the appendix	79269	Mucopolysaccharidosis type IIIA	1851	Multicystic renal dysplasia
		79270	Mucopolysaccharidosis type IIIB		
424053	Mucinous cystadenocarcinoma of pancreas	79271	Mucopolysaccharidosis type IIIC	48162	Multifocal acquired demyelinating sensory and motor neuropathy
		79272	Mucopolysaccharidosis type IIID	3282	Multifocal atrial tachycardia
319322	Mucinous tubular and spindle cell renal carcinoma	93474	Mucopolysaccharidosis type IS	99873	Multifocal eosinophilic granuloma
575	Muckle-Wells syndrome	582	Mucopolysaccharidosis type IV		
		309297	Mucopolysaccharidosis type IVA	464321	Multifocal lymphangioendotheliomatosis with thrombocytopenia
2331	Mucocutaneous lymph node syndrome	309310	Mucopolysaccharidosis type IVB		
		67041	Mucopolysaccharidosis type IX		
2451	Mucocutaneous venous malformations	583	Mucopolysaccharidosis type VI	464321	Multifocal lymphangioendotheliomatosis-thrombocytopenia syndrome
423461	Mucopolipidosis type 3 alpha/beta	276212	Mucopolysaccharidosis type VI, rapidly progressing	641	Multifocal motor neuropathy
423470	Mucopolipidosis type 3 gamma				
576	Mucopolipidosis type II				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
641	Multifocal motor neuropathy with conduction block	523	Multiple cutaneous and uterine leiomyomas	321	Multiple osteochondromas
2033	Multifocal muscular fibrosis-obstructed vessels syndrome	3453	Multiple endocrine deficiency-Addison disease-candidiasis syndrome	324299	Multiple paragangliomas associated with erythrocytosis
99003	Multifocal pattern dystrophy simulating fundus flavimaculatus			324299	Multiple paragangliomas associated with polycythemia
99003	Multifocal pattern dystrophy simulating Stargardt disease	652	Multiple endocrine neoplasia type 1	95494	Multiple pituitary hormone deficiencies, genetic forms
3286	Multifocal ventricular premature beats	653	Multiple endocrine neoplasia type 2	→1234	Multiple pterygium syndrome, Aslan type
		247698	Multiple endocrine neoplasia type 2A	2215	Multiple pterygium-malignant hyperthermia syndrome
319287	Multilocular clear cell adenocarcinoma	247709	Multiple endocrine neoplasia type 2B	3151	Multiple sclerosis-ichthyosis-factor VIII deficiency syndrome
319287	Multilocular clear cell carcinoma	247709	Multiple endocrine neoplasia type 3		
319287	Multilocular clear cell renal cell adenocarcinoma	276152	Multiple endocrine neoplasia type 4	65748	Multiple self-healing squamous epithelioma
319287	Multilocular clear cell renal cell carcinoma	166002	Multiple epiphyseal dysplasia due to collagen 9 anomaly	585	Multiple sulfatase deficiency
97366	Multilocular cyst of the kidney	93308	Multiple epiphyseal dysplasia type 1	2398	Multiple symmetric lipomatosis
		93307	Multiple epiphyseal dysplasia type 4	3237	Multiple synostoses syndrome
319287	Multilocular cystic clear cell renal cell neoplasm of low malignant potential	93311	Multiple epiphyseal dysplasia type 5	102	Multiple system atrophy
319287	Multilocular cystic renal cell adenocarcinoma	166016	Multiple epiphyseal dysplasia with Robin phenotype	227510	Multiple system atrophy, cerebellar type
		166024	Multiple epiphyseal dysplasia, Al-Gazali type	98933	Multiple system atrophy, parkinsonian type
319287	Multilocular cystic renal cell carcinoma	166011	Multiple epiphyseal dysplasia, Beighton type	102	Multisystem atrophy
168816	Multilocular peritoneal inclusion cyst	166016	Multiple epiphyseal dysplasia, Lowry type	404463	Multisystemic smooth muscle dysfunction syndrome
97366	Multilocular renal cyst	166032	Multiple epiphyseal dysplasia, with miniepiphyseal	2959	Mulvihill-Smith syndrome
97366	Multiloculated renal cyst	166029	Multiple epiphyseal dysplasia, with severe proximal femoral dysplasia	2578	MURCS association
598	Multiminicore disease			83315	Murine typhus
598	Multiminicore myopathy	166024	Multiple epiphyseal dysplasia-macrocephaly-distinctive facies syndrome	2028	Murray-Puretic-Drescher syndrome
2091	Multinodular goiter-cystic kidney-polydactyly syndrome	166011	Multiple epiphyseal dysplasia-myopia-deafness syndrome	99849	Muscle enolase deficiency
26791	Multiple acyl-CoA dehydrogenase deficiency	50920	Multiple fibroadenoma of the breast	171445	Muscle filaminopathy
394532	Multiple acyl-CoA dehydrogenase deficiency, mild type			97234	Muscle phosphoglycerate mutase deficiency
394529	Multiple acyl-CoA dehydrogenase deficiency, severe neonatal type	83454	Multiple glomus tumors	588	Muscle-eye-brain disease
2505	Multiple benign circumferential skin creases on limbs	201	Multiple hamartoma syndrome	370997	Muscle-eye-brain disease with bilateral multicystic leucodystrophy
		2300	Multiple intestinal atresia	588	Muscle-eye-brain syndrome
2678	Multiple café-au-lait spots	284139	Multiple joint dislocations-short stature-craniofacial dysmorphism-congenital heart defects syndrome	2576	Muscle-liver-brain-eye nanism
2678	Multiple café-au-lait syndrome	294049	Multiple joint dislocations-short stature-hyperlaxity-craniofacial dysmorphism syndrome	2579	Muscular atrophy-ataxia-retinitis pigmentosa-diabetes mellitus syndrome
321	Multiple cartilaginous exostoses			424261	Muscular dystrophy with progressive weakness, distal contractures and rigid spine
254519	Multiple congenital anomalies due to 14q32.2 maternally expressed gene defect	493	Multiple keratoacanthoma	199340	Muscular dystrophy, Selcen type
280633	Multiple congenital anomalies-hypotonia-seizures syndrome	65748	Multiple keratoacanthoma, Ferguson-Smith type	1877	Muscular dystrophy-white matter spongiosis syndrome
300496	Multiple congenital anomalies-hypotonia-seizures syndrome type 2	587	Multiple keratoacanthoma, Muir-Torre type	99849	Muscular enolase deficiency
1486	Multiple contracture syndrome, Finnish type	79455	Multiple mastocytoma	324416	Muscular hypertrophy-hepatomegaly-polyhydramnios syndrome
		29073	Multiple myeloma	2349	Muscular pseudohypertrophy-hypothyroidism syndrome
137776	Multiple contracture syndrome, Israeli-Bedouin type	→636	Multiple non-ossifying fibromatosis		
		435329	Multiple ossifying fibroma		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
494	Mutilating keratoderma of Vohwinkel	182050	MYH9-RD	289380	Myosclerosis
		182050	MYH9-related disease	337	Myositis ossificans progressiva
494	Mutilating keratoderma plus deafness	182050	MYH9-related disorder	764	Myositis purulenta tropica
		182050	MYH9-related syndrome	764	Myositis tropicans
659	Mutilating palmoplantar hyperkeratosis with periorificial keratotic plaques	182050	MYH9-related syndromic thrombocytopenia	306553	Myospherulosis
		2588	Myhre syndrome	614	Myotonia congenita
659	Mutilating palmoplantar keratoderma with periorificial keratotic plaques	109	Myhre-Riley-Smith syndrome	99734	Myotonia fluctuans
		480491	MYO5B-related progressive familial intrahepatic cholestasis	99735	Myotonia permanens
247798	MUTYH-related AFAP	45	Myoadenylate deaminase deficiency	3101	Myotonia-intellectual disability-skeletal anomalies syndrome
247798	MUTYH-related attenuated familial adenomatous polyposis			99736	Myotonia-painful contractions syndrome
247798	MUTYH-related attenuated familial polyposis coli	1942	Myoclonic atonic epilepsy	800	Myotonic chondrodystrophy
247798	MUTYH-related attenuated FAP	36899	Myoclonic dystonia	273	Myotonic dystrophy type 1
29	MVA	→36899	Myoclonic dystonia 15	606	Myotonic dystrophy type 2
2290	MVID	86913	Myoclonic epilepsy in non-progressive encephalopathies	→52430	Myotonic dystrophy type 3
2582	Myalgia-eosinophilia syndrome associated with tryptophan	86909	Myoclonic epilepsy of infancy	800	Myotonic myopathy, dwarfism, chondrodystrophy, ocular and facial anomalies
		1942	Myoclonic-astatic epilepsy		
589	Myasthenia gravis	1942	Myoclonic-astatic epilepsy in early childhood	79105	Myxofibrosarcoma
2583	Mycetoma	435438	Myoclonus epilepsy and ataxia due to potassium channel mutation	79105	Myxoid malignant fibrous histiocytoma
314946	Mycobacterium xenopi infection			99967	Myxoid/round cell liposarcoma
268249	Mycophenolate mofetil embryopathy	551	Myoclonus epilepsy associated with ragged-red fibres	57782	Myxoma with fibrous dysplasia
83482	Mycoplasma encephalitis	86913	Myoclonus epilepsy in non-progressive encephalopathies	1359	Myxoma-spotty pigmentation-endocrine overactivity syndrome
2584	Mycosis fungoides, Alibert-Bazin type			251643	Myxopapillary ependymoma
178512	Mycosis fungoides-associated follicular mucinosis	2589	Myoclonus-cerebellar ataxia-deafness syndrome	2608	N syndrome
		36899	Myoclonus-dystonia syndrome	79270	N-acetyl-alpha-glucosaminidase deficiency
183713	MyD88 deficiency	→36899	Myoclonus-dystonia type 15	583	N-acetylgalactosamine 4-sulfatase deficiency
59298	Myelinoclastic diffuse sclerosis	163696	Myoclonus-nephropathy syndrome		
135	Myelinosis centralis diffusa	178464	Myofibrillar myopathy with early respiratory failure	309297	N-acetylgalactosamine-6-sulfate sulfatase deficiency
2585	Myelocerebellar disorder			576	N-acetylglucosamine 1-phosphotransferase deficiency
268813	Myelocystocele	104077	Myopathic intestinal pseudoobstruction	79329	N-acetylglucosaminyltransferase 2 deficiency
86841	Myelodysplastic syndrome associated with isolated del(5q) chromosome abnormality	2596	Myopathy and diabetes mellitus		
		88635	Myopathy due to casequestrin and SERCA1 protein overload	137754	N-acyl-L-amino acid amidohydrolase deficiency
824	Myelofibrosis with myeloid metaplasia	97234	Myopathy due to phosphoglycerate mutase deficiency	103908	Na-H exchange deficiency
168953	Myeloid neoplasm associated with FGFR1 rearrangement	43115	Myopathy with exercise intolerance, Swedish type	178303	Nabius mask-like facial syndrome
168947	Myeloid neoplasm associated with PDGFRA rearrangement	171889	Myopathy with hexagonally cross-linked tubular arrays	439196	NAE
168950	Myeloid neoplasm associated with PDGFRB rearrangement			69087	Naegeli syndrome
86850	Myeloid sarcoma	2598	Myopathy, lactic acidosis and sideroblastic anemia	69087	Naegeli-Franceschetti-Jadassohn syndrome
29073	Myelomatosis	2601	Myopathy-growth delay-intellectual disability-hypospadias syndrome	840	Naevus syringocystadenomatosus papilliferus
93969	Myelomeningocele			245	NAFD
2587	Myeloperoxidase deficiency	1358	Myopathy-Moebius-Robin syndrome	3137	NAGA deficiency
437572	MYH7-related late-onset scapuloperoneal muscular dystrophy	289685	Myopericytoma	79279	NAGA deficiency type 1
		368	Myophosphorylase deficiency	79280	NAGA deficiency type 2
437572	MYH7-related late-onset scapuloperoneal syndrome	178493	Myopic macular degeneration	79281	NAGA deficiency type 3
437572	MYH7-related late-onset SPMD	178493	Myopic maculopathy	245	Nager acrofacial dysostosis
				245	Nager syndrome

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927	NAGS deficiency	157850	NBIA1	446	Neonatal hemochromatosis
2211	Naguib-Richieri-Costa syndrome	216873	NBIA1, atypical form	398097	Neonatal Hughes syndrome
423454	Nail and teeth abnormalities-marginal palmoplantar keratoderma-oral hyperpigmentation syndrome	216866	NBIA1, classic form	137577	Neonatal hypoxic and ischemic brain injury
		289560	NBIA4	59303	Neonatal ichthyosis-sclerosing cholangitis syndrome
		329284	NBIA5	294023	Neonatal inflammatory skin and bowel disease
		397725	NBIA6	247598	Neonatal intrahepatic cholestasis caused by citrin deficiency
→1487	Nail dysplasia-camptodactyly-brachydactyly type B syndrome	289560	NBIA due to C19orf12 mutation	247598	Neonatal intrahepatic cholestasis due to citrin deficiency
		647	NBS	238688	Neonatal iodine exposure
2614	Nail-patella syndrome	240760	NBS-like disorder	398124	Neonatal lupus erythematosus
2613	Nail-patella-like renal disease	240760	NBSLD	284979	Neonatal Marfan syndrome
158676	Nails-only DEB	217560	NCHI	69063	Neonatal membranous glomerulopathy with maternal NEP deficiency
853	NAIT	1947	NCL, Northern epilepsy variant	69063	Neonatal membranous glomerulopathy with maternal neutral endopeptidase deficiency
101	Naito-Oyanagi disease	2481	NCM	284979	Neonatal MFS
2229	Najjar syndrome	75327	NCMD	79242	Neonatal multiple carboxylase deficiency
1063	Nakagawa angioblastoma	91495	NCRNA disease	391504	Neonatal myasthenia gravis
2615	Nakajo-Nishimura syndrome	443162	NDE1-related microhydranencephaly	→42738	Neonatal neutropenia
2822	Nakamura-Osame syndrome	399103	Nebulin-related early-onset distal myopathy	289857	Neonatal NKH
44	NALD	158011	Necrobiotic xanthogranuloma	289857	Neonatal non-ketotic hyperglycemia
206569	NAM	439196	Necrolytic acral erythema	56304	Neonatal osseous dysplasia type 1
→1359	NAME syndrome	391673	Necrotizing enterocolitis	3455	Neonatal progeroid syndrome
383	Nance deafness	440368	Necrotizing soft tissue infection	70587	Neonatal respiratory distress syndrome
627	Nance-Horan syndrome	217560	NEHI	3206	Neonatal Schwartz-Jampel syndrome
35612	Nanophthalmia	464366	NEK9-related lethal skeletal dysplasia	398127	Neonatal scleroderma
85196	NAO syndrome	199244	Nelson syndrome	466784	Neonatal severe cardiopulmonary failure due to mitochondrial methylation defect
247868	NAPS12	217563	Neonatal acute respiratory distress due to SP-B deficiency	417	Neonatal severe primary hyperparathyroidism
83465	Narcolepsy without cataplexy	217563	Neonatal acute respiratory distress due to surfactant protein B deficiency	1451	Neonatal-onset multisystem inflammatory disease
2073	Narcolepsy-cataplexy syndrome	44	Neonatal adrenoleukodystrophy	314950	Neoplastic hypereosinophilic syndrome
644	NARP syndrome	398109	Neonatal AHA	94058	Neovascular glaucoma
141103	Nasal dermoid cyst	398109	Neonatal AIHA	654	Nephroblastoma
141103	Nasal dermoid sinus cyst	464370	Neonatal alloimmune neutropenia	2849	Nephroblastomatosis-fetal ascites-macrosomia-Wilms tumor syndrome
141219	Nasal dorsum fistula/cyst	398097	Neonatal antiphospholipid antibody syndrome	223	Nephrogenic diabetes insipidus
141118	Nasal encephalocele	398097	Neonatal antiphospholipid syndrome	3145	Nephrogenic diabetes insipidus-intracranial calcification syndrome
141115	Nasal ganglioglioma	398109	Neonatal autoimmune hemolytic anemia	137617	Nephrogenic fibrosing dermatopathy
141112	Nasal glial heterotopia	137929	Neonatal brainstem dysfunction	93606	Nephrogenic syndrome of inappropriate antidiuresis
141112	Nasal glioma	314911	Neonatal Canavan disease		
86879	Nasal T/natural killer-cell lymphoma	313906	Neonatal congenital pancreatic cyst		
2662	Nasodigitoacoustic syndrome	398117	Neonatal dermatomyositis		
141083	Nasolacrimal duct cyst	79118	Neonatal diabetes-congenital hypothyroidism-congenital glaucoma-hepatic fibrosis-polycystic kidneys syndrome		
141083	Nasolacrimal mucocele	398117	Neonatal DM		
2399	Nasopalpebral lipoma-coloboma syndrome	457185	Neonatal encephalomyopathy-cardiomyopathy-respiratory distress syndrome		
150	Nasopharyngeal carcinoma	289857	Neonatal glycine encephalopathy		
141107	Nasopharyngeal teratoma				
2770	Nasu-Hakola disease				
1654	Natal teeth-intestinal pseudoobstruction-patent ductus syndrome				
2663	Nathalie syndrome				
168572	Native American myopathy				
69739	Navajo brainstem syndrome				
255229	Navajo neurohepatopathy				
255229	Navajo neuropathy				
34217	Naxos disease				
377	NBCCS				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
137617	Nephrogenic systemic fibrosis	157850	Neurodegeneration with brain iron accumulation type 1	637	Neurofibromatosis type 2
93622	Nephrolithiasis type 1			93921	Neurofibromatosis type 3
93623	Nephrolithiasis type 2	216873	Neurodegeneration with brain iron accumulation type 1, atypical form	2678	Neurofibromatosis type 6
655	Nephronophthisis			638	Neurofibromatosis-Noonan syndrome
3156	Nephronophthisis with retinal dystrophy	216866	Neurodegeneration with brain iron accumulation type 1, classic form	3148	Neurofibrosarcoma
84081	Nephronophthisis-hepatic fibrosis syndrome	289560	Neurodegeneration with brain iron accumulation type 4	970	Neurogenic acroosteolysis
411629	Nephropathic infantile cystinosis	329284	Neurodegeneration with brain iron accumulation type 5	1143	Neurogenic arthrogryposis multiplex congenita
2668	Nephropathy-deafness-hyperparathyroidism syndrome			100073	Neurogenic cervical rib syndrome
2669	Nephrosis-deafness-urinary tract-digital malformations syndrome	217382	Neurodegenerative syndrome due to cerebral folate transport deficiency	100073	Neurogenic costoclavicular syndrome
2065	Nephrosis-neuronal dysmigration syndrome	453499	Neurodevelopmental disorder-craniofacial dysmorphism-cardiac defect-hip dysplasia syndrome	178029	Neurogenic diabetes insipidus
300333	Nephrotic syndrome-deafness-pretibial epidermolysis bullosa syndrome			644	Neurogenic muscle weakness-ataxia-retinitis pigmentosa syndrome
300333	Nephrotic syndrome-hearing loss-pretibial epidermolysis bullosa syndrome	352665	Neurodevelopmental disorder-craniofacial dysmorphism-cardiac defect-hip dysplasia syndrome due to 9q21 microdeletion	98593	Neurogenic palpebral tumor
2337	NEPPK			3148	Neurogenic sarcoma
280576	Nestor-Guillermo progeria syndrome	453504	Neurodevelopmental disorder-craniofacial dysmorphism-cardiac defect-hip dysplasia syndrome due to a point mutation	431255	Neurogenic scapulo-peroneal amyotrophy, New England type
100082	NET of the anal canal			85146	Neurogenic scapulo-peroneal syndrome, Kaeser type
100080	NET of the colon	3474	Neuroectodermal dysplasia, CHIME type	100073	Neurogenic thoracic outlet compression syndrome
634	Netherton syndrome	33445	Neuroectodermal melanolyosomal disease	100073	Neurogenic thoracic outlet syndrome
2671	Neu-Laxova syndrome	3474	Neuroectodermal syndrome, Zurich type	100073	Neurogenic TOS
99078	Neuhauser anomaly			94093	Neuroleptic malignant syndrome
2479	Neuhäuser syndrome	2676	Neuroectodermal-endocrine syndrome	36397	Neurolipomatosis
3350	Neuhauser-Daly-Magnelli syndrome			163746	Neurologic Waardenburg-Shah syndrome
2672	Neuhauser-Eichner-Opitz syndrome	217560	Neuroendocrine cell hyperplasia of infancy	137754	Neurological conditions associated with aminoacylase 1 deficiency
2901	Neuralgic amyotrophy			206586	Neurolymphomatosis
2901	Neuralgic shoulder amyotrophy	100082	Neuroendocrine neoplasm of the anal canal	71211	Neuromyelitis optica
351	Neuraminidase deficiency with beta-galactosidase deficiency	100080	Neuroendocrine neoplasm of the colon	1947	Neuronal ceroid lipofuscinosis, Northern epilepsy variant
268865	Neurenteric cyst	100081	Neuroendocrine tumor of rectum	99811	Neuronal intestinal pseudoobstruction
252164	Neurilemmoma	100082	Neuroendocrine tumor of the anal canal		
93921	Neurilemmomatosis	100079	Neuroendocrine tumor of the appendix	2289	Neuronal intranuclear inclusion disease
252164	Neurilemoma			639	Neuropathy associated with monoclonal IgM antibodies to myelin-associated glycoprotein
635	Neuroblastoma	100080	Neuroendocrine tumor of the colon	139512	Neuropathy with hearing impairment
2481	Neurocutaneous melanocytosis	2677	Neuroepithelioma		
2481	Neurocutaneous melanosis	2673	Neurofaciodigitorenal syndrome	644	Neuropathy-ataxia-retinitis pigmentosa syndrome
35664	Neurocutaneous syndrome, Bicknell type	157846	Neuroferritinopathy		
88639	Neurodegeneration due to 3-hydroxyisobutyryl-CoA hydrolase deficiency	252183	Neurofibroma	217622	Neurosensory deafness with dilated cardiomyopathy
		137605	Neurofibromatosis 1-like syndrome		
		636	Neurofibromatosis type 1	217622	Neurosensory hearing loss with dilated cardiomyopathy
289560	Neurodegeneration with brain iron accumulation due to C19orf12 mutation	363700	Neurofibromatosis type 1 due to NF1 mutation or intragenic deletion	137596	Neurotrophic keratitis
397725	Neurodegeneration with brain iron accumulation due to COASY mutation	97685	Neurofibromatosis type 1 microdeletion syndrome	137596	Neurotrophic keratopathy
		638	Neurofibromatosis type 1-Noonan syndrome		

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98907	Neutral lipid storage disease with ichthyosis	141179	NICH	436166	NLRC4-related macrophage activation syndrome
		3051	Nicolaides-Baraitser syndrome		
98908	Neutral lipid storage disease with myopathy without ichthyosis	77292	Niemann-Pick disease type A	436166	NLRC4-related MAS
		77293	Niemann-Pick disease type B	247868	NLRP12-associated hereditary periodic fever syndrome
98908	Neutral lipid storage myopathy	646	Niemann-Pick disease type C		
→86872	Neutropenia-hyperlymphocytosis with large granular lymphocytes syndrome	216986	Niemann-Pick disease type C, adult neurologic onset	98907	NLSDI
				98908	NLSDM
2690	Neutropenia-monocytopenia-deafness syndrome	216981	Niemann-Pick disease type C, classic form	443167	NMC
				391504	NMG
183707	Neutrophil immunodeficiency syndrome	216981	Niemann-Pick disease type C, juvenile neurologic onset	86867	NMZL
				2615	NNS
169142	Neutrophil-specific granule deficiency	216978	Niemann-Pick disease type C, late infantile neurologic onset	1884	Noble-Bass-Sherman syndrome
				31204	Nocardiosis
575	Neutrophilic urticaria	216975	Niemann-Pick disease type C, severe early infantile neurologic onset	→98784	Nocturnal paroxysmal dystonia
370059	NEVADA syndrome			86867	Nodal marginal zone B-cell lymphoma
→1359	Nevi-atrial myxoma-myxoid neurofibromata-ephelides syndrome	216972	Niemann-Pick disease type C, severe perinatal form	137810	Nodular cutaneous amyloidosis
→1900	Nevo syndrome	→646	Niemann-Pick disease type D	477742	Nodular fasciitis
		99022	Niemann-Pick disease type E	90393	Nodular lichen myxedematosus
377	Nevoid basal cell carcinoma syndrome	→646	Niemann-Pick disease, Nova Scotia type	86893	Nodular lymphocyte predominant Hodgkin lymphoma
228264	Nevus anelasticus	2633	Nievergelt syndrome	2149	Nodular neuronal heterotopia
64754	Nevus comedonicus syndrome			33577	Nodular non-suppurative panniculitis
228254	Nevus elasticus	1390	Night blindness-skeletal anomalies-dysmorphism syndrome		
370059	Nevus epidermicus verrucosus with angiodysplasia and aneurysms	98757	Nigro-spino-dentatal degeneration with nuclear ophthalmoplegia	48372	Nodular regenerative hyperplasia of the liver
263432	Nevus fuscocaeruleus acromiodeltoideus	432	nIHH	158772	Nodular urticaria pigmentosa
		2322	Niikawa-Kuroki syndrome	85196	Nodulosis-arthropathy-osteolysis syndrome
263425	Nevus fuscoceruleus ophthalmomaxillaris	647	Nijmegen breakage syndrome	2700	Noma
263432	Nevus of Ito	240760	Nijmegen breakage syndrome-like disorder	1451	NOMID syndrome
263425	Nevus of Ota	447731	NIK deficiency	73267	Non-24-hour sleep-wake syndrome
2612	Nevus sebaceus of Jadassohn	781	Nine Mile fever		
2612	Nevus sebaceus syndrome	99825	Nipah encephalitis	231720	Non-acquired combined pituitary hormone deficiency with spine abnormalities
363558	New-onset refractory status epilepticus	99825	Nipah fever		
		99825	Nipah virus disease	631	Non-acquired isolated growth hormone deficiency
83471	Nezelof syndrome	59303	NISCH syndrome		
636	NF1	1422	Nivelon-Nivelon-Mabille syndrome	97566	Non-amyloid fibrillary glomerulonephritis
97685	NF1 microdeletion syndrome	263665	NK-cell enteropathy		
137605	NF1-like syndrome	86873	NK-cell large granular lymphocyte leukemia	97566	Non-amyloid fibrillary glomerulopathy
637	NF2			86861	Non-amyloid MIDD
93921	NF3	86873	NK-cell LGL leukemia		
2678	NF6	86879	NK/T-cell lymphoma	86861	Non-amyloid monoclonal immunoglobulin deposition disease
69087	NFJ syndrome	407	NKA		
638	NFNS	86879	NKTCL	79394	Non-bullous congenital ichthyosiform erythroderma
91349	NFPA	86893	NLPHL		
401869	NFU1 deficiency			289362	Non-central nervous system-localized embryonal carcinoma
289356	NGCO	436166	NLRC4-related autoinflammatory syndrome with macrophage activation syndrome		
404454	NGLY1 deficiency			77259	Non-cerebral juvenile Gaucher disease
404454	NGLY1-CDDG	436166	NLRC4-related autoinflammatory syndrome with MAS	48372	Non-cirrhotic nodulation
280576	NGPS			854	Non-cirrhotic portal vein thrombosis
2770	NHD			325529	Non-classic congenital lipid adrenal hyperplasia due to STAR deficiency
169079	NHEJ1 deficiency	436166	NLRC4-related infantile enterocolitis-autoinflammatory syndrome		
276608	NI-PHH				
247598	NICCD				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
289362	Non-CNS-localized embryonal carcinoma	329918	Non-immunoglobulin-mediated MPGN	35093	Non-syndromic sagittal synostosis
216796	Non-deforming osteogenesis imperfecta	263548	Non-inflammatory generalized peeling skin syndrome type A.	35098	Non-syndromic unicoronal synostosis
96136	Non-distal deletion 7p	263548	Non-inflammatory peeling skin syndrome type A	96136	Non-telomeric monosomy 7p
1581	Non-distal deletion 10q			1581	Non-telomeric monosomy 10q
96160	Non-distal deletion 12q	141179	Non-involuting congenital hemangioma	96160	Non-telomeric monosomy 12q
96164	Non-distal deletion 20q			96164	Non-telomeric monosomy 20q
96112	Non-distal duplication 9q	407	Non-ketotic hyperglycinemia	3306	Non-telomeric tetrasomy 15q
1695	Non-distal duplication 10q	98890	Non-Leber type optic atrophy with early-onset	96112	Non-telomeric trisomy 9q
1702	Non-distal duplication 13q			1695	Non-telomeric trisomy 10q
96136	Non-distal monosomy 7p	411641	Non-nephropathic cystinosis	1702	Non-telomeric trisomy 13q
1581	Non-distal monosomy 10q	84085	Non-neurogenic neurogenic bladder	411703	Non-tuberculous mycobacterial lung disease
96160	Non-distal monosomy 12q	209989	Non-papillary transitional cell carcinoma of the bladder	209919	Non-Wilsonian hepatic copper toxicosis of infancy and childhood
96164	Non-distal monosomy 20q	209989	Non-papillary urothelial carcinoma	602	Nonaka myopathy
3306	Non-distal tetrasomy 15q	238583	Non-phenylketonuric hyperphenylalaninemia	79452	Nonne-Milroy lymphedema
96112	Non-distal trisomy 9q	79651	Non-PKU HPA	648	Noonan syndrome
1695	Non-distal trisomy 10q	→144	Non-polyposis Turcot syndrome	500	Noonan syndrome with multiple lentigines
1702	Non-distal trisomy 13q	314647	Non-progressive cerebellar ataxia with intellectual disability	363972	Noonan syndrome-like disorder with JMML
329469	Non-DS-AMKL	1766	Non-progressive cerebellar ataxia-intellectual disability syndrome	363972	Noonan syndrome-like disorder with juvenile myelomonocytic leukemia
206538	Non-dysgerminomatous germ cell cancer of ovary			363972	Noonan syndrome-like disorder with loose anagen hair
363494	Non-dysgerminomatous germ cell tumor of testis	436271	Non-progressive predominantly posterior cavitating leukoencephalopathy with peripheral neuropathy	2701	Noonan syndrome-like disorder with loose anagen hair
2337	Non-epidermolytic palmoplantar keratoderma			230	Noradrenaline deficiency
→2199	Non-epidermolytic palmoplantar keratoderma	439202	Non-recovering OBPI	230	Norepinephrine deficiency
2972	Non-eruption of teeth-maxillary hypoplasia-genu valgum syndrome	439202	Non-recovering OBPL	314928	Normal pressure hydrocephalus
100070	Non-fluent variant PPA	439202	Non-recovering obstetric brachial plexus lesion	2254	Norman disease
91349	Non-functioning pituitary adenoma			79255	Norman-Landing disease
26137	Non-giant cell granulomatous temporal arteritis with eosinophilia	101106	Non-secreting chemodectoma	→682	Normokalemic periodic paralysis
→79452	Non-hereditary congenital primary lymphedema	94080	Non-secreting paraganglioma	→682	Normokalemic PP
→90186	Non-hereditary late-onset primary lymphedema	363494	Non-seminomatous germ cell tumor of testis	→682	NormoKPP
357034	Non-hereditary retinoblastoma	91364	Non-specific idiopathic interstitial pneumonia	812	Normomorphic sialidosis
163924	Non-herpetic acute limbic encephalitis	91364	Non-specific interstitial pneumonia	→682	NormoPP
329883	Non-hypoproteinemic hypertrophic gastropathy	206572	Non-specific myositis	432	Normosmic congenital hypogonadotropic hypogonadism
329918	Non-Ig-mediated membranoproliferative glomerulonephritis	90031	Non-spherocytic hemolytic anemia due to hexokinase deficiency	432	Normosmic idiopathic hypogonadotropic hypogonadism
329918	Non-Ig-mediated MPGN	35099	Non-syndromic bicoronal synostosis	649	Norrie disease
363999	Non-immune fetal edema	30391	Non-syndromic biliary atresia	649	Norrie-Warburg disease
363999	Non-immune fetal hydrops	91495	Non-syndromic congenital retinal non-attachment	363558	NORSE
363999	Non-immune HF	49042	Non-syndromic dentinogenesis imperfecta	75327	North Carolina macular dystrophy
363999	Non-immune hydrops fetalis	49042	Non-syndromic DGI	75327	North Carolina macular dystrophy, retinal 1
329918	Non-immunoglobulin-mediated membranoproliferative glomerulonephritis	276234	Non-syndromic male infertility due to sperm motility disorder	280620	North Sea progressive myoclonus epilepsy
		3366	Non-syndromic metopic craniosynostosis	1947	Northern epilepsy
				79293	Norum disease
				1134	Nose agenesis
				178	Notochordal sarcoma
				2703	Nova syndrome
				2005	Novak syndrome

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
314928	NPH	217017	Occipital atretic cephalocele-unusual facies-large feet syndrome	2720	Oculocerebral hypopigmentation syndrome, Preus type
3032	NPHP3-related Meckel-like syndrome	268823	Occipital encephalocele	1647	Oculocerebrocutaneous syndrome
480476	NR1H4 deficiency	198	Occipital horn syndrome	2707	Oculocerebrofacial syndrome, Kaufman type
634	NS	280640	Occipital malformations of cortical development	534	Oculocerebrorenal dystrophy
88616	NS-ARID	280640	Occipital MCD	534	Oculocerebrorenal syndrome of Lowe
2701	NS/LAH	280640	Occipital pachygyria and polymicrogyria	352731	Oculocutaneous albinism type 1
417	NSHPT	353351	Occlusive idiopathic juxtafoveal retinal telangiectasis	79431	Oculocutaneous albinism type 1A
93606	NSIAD	51608	Occlusive infantile arteriopathy	79432	Oculocutaneous albinism type 1B
91364	NSIP	1647	OCCS	79433	Oculocutaneous albinism type 2
454840	NTHL1-related AFAP	99889	Occult ectopic ACTH secretion	79435	Oculocutaneous albinism type 3
454840	NTHL1-related attenuated familial adenomatous polyposis	247834	Occult macular dystrophy	370091	Oculocutaneous albinism type 4
100073	NTOS	84085	Occult neuropathic bladder	370097	Oculocutaneous albinism type 5
314790	Null pituitary adenoma	2704	Ochoa syndrome	352745	Oculocutaneous albinism type 6
280234	Null syndrome	247834	OCMD	79434	Oculocutaneous albinism type 7
443167	NUT midline carcinoma	534	OCR	28378	Oculocutaneous albinism, Amish type
54	OA1	534	OCRL	2709	Oculocutaneous tyrosinemia
398156	OAFNS	664	OCT deficiency	2710	Oculodental syndrome, Rutherford type
1106	OAS	54	Ocular albinism type 1	2710	Oculodentodigital dysplasia
374	OAV dysplasia	352740	Ocular albinism with congenital sensorineural deafness	2710	Oculodentoosseous dysplasia
374	OAVS	1000	Ocular albinism with late-onset sensorineural deafness	3339	Oculoectodermal syndrome
97297	Oberklaid-Danks syndrome	54	Ocular albinism, Nettleship-Falls type	2712	Oculofaciocardidental syndrome
397615	Obesity due to CEP19 deficiency	411641	Ocular cystinosis	1876	Oculogastrointestinal muscular dystrophy
66628	Obesity due to congenital leptin deficiency	2788	Ocular form of osteogenesis imperfecta	2108	Oculomandibulofacial syndrome
179494	Obesity due to leptin receptor gene deficiency	1125	Ocular motor apraxia, Cogan type	1794	Oculomaxillofacial dysostosis
71529	Obesity due to melanocortin 4 receptor deficiency	99922	Ocular pemphigoid	1154	Oculomelic amyoplasia
71526	Obesity due to pro-opiomelanocortin deficiency	534	Oculo-cerebro-renal dystrophy	1125	Oculomotor apraxia, Cogan type
71528	Obesity due to prohormone convertase I deficiency	534	Oculo-cerebro-renal syndrome	2713	Oculoosteocutaneous syndrome
369873	Obesity due to SIM1 deficiency	1305	Oculo-digito-esophageal-duodenal syndrome	99806	Oculootodontal syndrome
88643	Obesity-colitis-hypothyroidism-cardiac hypertrophy-developmental delay syndrome	391641	Oculo-digito-esophageal-duodenal syndrome type 1	→293843	Oculopalatoskeletal syndrome
1303	Obliterative bronchiolitis	→1200	Oculo-oto-facial dysplasia	98897	Oculopharyngeal distal myopathy
64743	Obliterative portal venopathy	2307	Oculo-oto-radial syndrome	270	Oculopharyngeal muscular dystrophy
2970	Obrinsky syndrome	2714	Oculo-palato-cerebral dwarfism	98897	Oculopharyngodistal myopathy
3411	Obstructed hemivagina and ipsilateral renal anomaly	2714	Oculo-palato-cerebral syndrome	2715	Oculorenocerebellar syndrome
352731	OCA1	→293843	Oculo-skeletal-abdominal syndrome	2717	Oculotrichoanal syndrome
352734	OCA1-MP	2716	Oculo-skeletal-renal syndrome	2718	Oculotrichodysplasia
352737	OCA1-TS	157962	Oculoauricular syndrome, Schorderet type	166272	ODCD
79431	OCA1A	398156	Oculoauriculofrontonasal syndrome	2710	ODDD syndrome
79434	OCA1B	374	Oculoauriculovertebral dysplasia	1305	ODED syndrome
79432	OCA2	2549	Oculoauriculovertebral spectrum with radial defects	391641	ODED syndrome type 1
79433	OCA3	374	Oculoauriculovertebral syndrome	999	O'Doherty syndrome
79435	OCA4	2705	Oculocerebral dysplasia	2253	O'Donnell-Pappas syndrome
370091	OCA5	2719	Oculocerebral hypopigmentation syndrome, Cross type	2722	Odonto-onycho dysplasia-alopecia syndrome
370097	OCA6			2721	Odonto-onycho-dermal dysplasia
352745	OCA7			→2036	Odonto-onycho-hypohidrotic dysplasia-midline scalp defects syndrome

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
69082	Odonto-tricho-ungual-digito-palmar syndrome	261638	Okhiro syndrome due to 20q13 microdeletion	210115	OMPP
				1183	OMS
69082	Odonto-tricho-ungual-digito-palmar syndrome, Mendoza-Valiente type	261647	Okhiro syndrome due to a point mutation	319266	Omsk hemorrhagic fever
166272	Odontochondrodysplasia	261638	Okhiro syndrome due to del(20)(q13)	3191	Onat syndrome
447777	Odontogenic keratocystoma			2737	Onchocerciasis
247685	Odontohypophosphatasia	261638	Okhiro syndrome due to monosomy 20q13	137675	Oncocytic cardiomyopathy
77295	Odontoleukodystrophy			352540	Oncogenic hypophosphatemic osteomalacia
2724	Odontomatosis-aortae esophagus stenosis syndrome	69088	OL-EDA-ID	352540	Oncogenic osteomalacia
		→113	Oley syndrome	661	Ondine curse
1811	Odontomicronychial dysplasia	478	Olfacto-genital pathological sequence	661	Ondine syndrome
2723	Odontotrichomelic syndrome			99803	Ondine-Hirschsprung disease
1487	ODP	1957	Olfactory neuroblastoma	99803	Ondine-Hirschsprung syndrome
93929	OEIS complex	85410	Oligoarticular JIA	→33364	ONMR syndrome
2676	Oerter-Friedman-Anderson syndrome	247839	Oligoarticular JIA with anti-nuclear antibodies	238744	Onycho-digito-mammary syndrome
2792	OFC syndrome	247846	Oligoarticular JIA without anti-nuclear antibodies	→33364	Onycho-tricho-dysplasia-neutropenia syndrome
2712	OFCD syndrome			300504	Onychocytic matricoma
2750	OFD1	85410	Oligoarticular juvenile idiopathic arthritis	79153	Onychodystrophy totalis
2751	OFD2			300512	Onychomatricoma
2752	OFD3	247839	Oligoarticular juvenile idiopathic arthritis with anti-nuclear antibodies	2614	Onychoosteodysplasia
2753	OFD4			2786	OCHS
2919	OFD5	247846	Oligoarticular juvenile idiopathic arthritis without anti-nuclear antibodies	99806	OOD
2754	OFD6			2721	OODD
→2750	OFD7	251656	Oligoastrocytoma	98890	OPA2
2755	OFD8			67036	OPA3, autosomal dominant
141007	OFD9	75378	Oligocone syndrome	49042	Opalescent teeth without OI
2756	OFD10	75378	Oligocone trichromacy	49042	Opalescent teeth without osteogenesis imperfecta
141000	OFD11	251627	Oligodendroglioma		
141327	OFD12	99798	Oligodontia	90650	OPD I syndrome
141330	OFD13	300576	Oligodontia-cancer predisposition syndrome	90652	OPD II syndrome
434179	OFD14	2260	Oligomeganephronia	90650	OPD syndrome 1
2750	OFDI			90652	OPD syndrome 2
2750	OFDSI	2260	Oligomeganephronic renal hypoplasia	98897	OPDM
391655	Off-periods in Parkinson disease not responding to oral treatment			137831	Oligophrenin-1 syndrome
424080	OGCT of pancreas	2920	Oliver syndrome	137831	OPHN1 syndrome
276432	Ogden syndrome	3363	Oliver-McFarlane syndrome	1106	Ophthalmoacromelic syndrome
75382	Oguchi disease	2732	Olivopontocerebellar atrophy-deafness syndrome	2741	Ophthalmomandibulomelic dysplasia
75382	Oguchi syndrome			1186	Ophthalmoplegia-hypotonia-ataxia-hypoacusis-athetosis syndrome
1186	Ohaha syndrome	296	Ollier disease	2743	Ophthalmoplegia-intellectual disability-lingua scrotalis syndrome
2728	Ohdo syndrome	659	Olmsted syndrome		
2728	Ohdo-Madokoro-Sonoda syndrome	1183	OMA syndrome	1308	Opitz C trigonocephaly
64739	OHSS	247834	OMD	2745	Opitz G/BBB syndrome
1934	Ohtahara syndrome	39041	Omenn syndrome	2745	Opitz syndrome
3411	OHVIRA syndrome	2741	OMM syndrome	1308	Opitz trigonocephaly C syndrome
666	OI	2733	Omodysplasia	1308	Opitz trigonocephaly syndrome
216796	OI type 1	660	Omphalocele	97297	Opitz trigonocephaly-like syndrome
216804	OI type 2	3164	Omphalocele syndrome, Shprintzen-Goldberg type	1786	Opitz-Caltabiano syndrome
216812	OI type 3			2745	Opitz-Frias syndrome
216820	OI type 4	93929	Omphalocele-cloacal exstrophy-imperforate anus-spinal defect syndrome	93932	Opitz-Kaveggia syndrome
216828	OI type 5			270	OPMD
2729	Okamoto syndrome	490	Omphalomesenteric cyst	256	Oppenheim dystonia
93293	Okhiro syndrome				

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2788	OPPG	415	Ornithine carrier deficiency	3314	Osteochondritis of phalangeal epiphyses
2746	Opsismodysplasia	664	Ornithine transcarbamylase deficiency	2054	Osteochondritis of tarsal/metatarsal bone
1183	Opsoclonus-myoclonus syndrome	415	Ornithine translocase deficiency	2380	Osteochondritis of the capital femoral epiphysis
1183	Opsoclonus-myoclonus-ataxia syndrome	415	ORNT1 deficiency	97332	Osteochondritis of the lunate bone
363746	Optic ataxia-gaze apraxia-simultanagnosia syndrome	2319	Orocraniodigital syndrome	97335	Osteochondritis of the tibial tubercle
98673	Optic atrophy type 1	353253	Orodynia	2653	Osteochondrodysplastic dwarfism-deafness-retinitis pigmentosa syndrome
98890	Optic atrophy type 2	2750	Orofaciodigital syndrome type 1	2653	Osteochondrodysplastic nanism-deafness-retinitis pigmentosa syndrome
1215	Optic atrophy-deafness-polyneuropathy-myopathy syndrome	2751	Orofaciodigital syndrome type 2	800	Osteochondromuscular dystrophy
401777	Optic atrophy-intellectual disability syndrome	2752	Orofaciodigital syndrome type 3	2768	Osteochondrosis deformans tibiae
→1215	Optic atrophy-ophthalmoplegia-ptosis-deafness-myopathy syndrome	2753	Orofaciodigital syndrome type 4	97337	Osteochondrosis of patella
313800	Optic nerve edema-splenomegaly syndrome	2919	Orofaciodigital syndrome type 5	3314	Osteochondrosis of phalangeal epiphyses
2086	Optic pathway glioma	2754	Orofaciodigital syndrome type 6	2380	Osteochondrosis of the capital femoral epiphysis
353253	Oral dysesthesia	→2750	Orofaciodigital syndrome type 7	97336	Osteochondrosis of the capital humerus
31142	Oral erosive lichen	2755	Orofaciodigital syndrome type 8	97332	Osteochondrosis of the lunate bone
357154	Oral submucous fibrosis	141007	Orofaciodigital syndrome type 9	2054	Osteochondrosis of the tarsal bone
457252	Oral tongue squamous cell carcinoma	2756	Orofaciodigital syndrome type 10	97335	Osteochondrosis of the tibial tubercle
2750	Oral-facial-digital syndrome type 1	141000	Orofaciodigital syndrome type 11	424080	Osteoclastic giant cell tumor of pancreas
2751	Oral-facial-digital syndrome type 2	141327	Orofaciodigital syndrome type 12	2763	Osteocraniosplenic syndrome
2752	Oral-facial-digital syndrome type 3	141330	Orofaciodigital syndrome type 13	2763	Osteocraniosplenosis
2753	Oral-facial-digital syndrome type 4	434179	Orofaciodigital syndrome type 14	666	Osteogenesis imperfecta
2919	Oral-facial-digital syndrome type 5	2756	Orofaciodigital syndrome with fibular aplasia	216796	Osteogenesis imperfecta type 1
2754	Oral-facial-digital syndrome type 6	141007	Orofaciodigital syndrome with retinal abnormalities	216804	Osteogenesis imperfecta type 2
→2750	Oral-facial-digital syndrome type 7	2755	Orofaciodigital syndrome, Edwards type	216812	Osteogenesis imperfecta type 3
2755	Oral-facial-digital syndrome type 8	141000	Orofaciodigital syndrome, Gabrielli type	216820	Osteogenesis imperfecta type 4
141007	Oral-facial-digital syndrome type 9	2919	Orofaciodigital syndrome, Thurston type	216828	Osteogenesis imperfecta type 5
2756	Oral-facial-digital syndrome type 10	93958	Oromandibular dystonia	2771	Osteogenesis imperfecta-congenital joint contractures syndrome
141000	Oral-facial-digital syndrome type 11	141077	Oropharyngeal teratoma	2773	Osteogenesis imperfecta-retinopathy-seizures-intellectual disability syndrome
141327	Oral-facial-digital syndrome type 12	30	Oroticaciduria	668	Osteogenic sarcoma
141330	Oral-facial-digital syndrome type 13	30	Orotidylic decarboxylase deficiency	2645	Osteoglophonic dwarfism
434179	Oral-facial-digital syndrome type 14	64692	Oroya fever	2777	Osteomesopyknosis
141007	Oral-facial-digital syndrome with retinal abnormalities	443236	Orthostatic intolerance due to NET deficiency	824	Osteomyelofibrosis
2755	Oral-facial-digital syndrome, Edwards type	→293843	OSA syndrome	399293	Osteonecrosis of the jaw
141000	Oral-facial-digital syndrome, Gabrielli type	93382	Osebold-Remondini syndrome	2780	Osteopathia striata-cranial sclerosis syndrome
1647	Orbital cyst with cerebral and focal dermal malformations	97335	Osgood-Schlatter disease	2779	Osteopathia striata-pigmentary dermopathy-white forelock syndrome
52994	Orbital leiomyoma	2760	OSLAM syndrome	2324	Osteopenia-intellectual disability-sparse hair syndrome
268139	Orbital medulloepithelioma	729	Osler-Vaquez disease		
2612	Organoid nevus syndrome	1427	OSMED		
166421	Orgasm-induced seizures	357154	OSMF		
49041	Ormond disease	140436	Osseous venous malformation		
414	Ornithine aminotransferase deficiency	73230	Ossification anomalies-psychomotor developmental delay syndrome		
664	Ornithine carbamoyltransferase deficiency	57196	Osteitis condensans of the clavicle		
		58040	Osteoblastoma		
		2764	Osteochondritis dissecans		
		251262	Osteochondritis dissecans and short stature		

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91133	Osteopenia-myopia-hearing loss-intellectual disability-facial dysmorphism syndrome	213512	Ovarian malignant mixed epithelial mesenchymal tumor	99736	Painful congenital myotonia
53	Osteopetrosis autosomal dominant type 2	213512	Ovarian malignant mixed Müllerian tumor	99736	Painful myotonia
2785	Osteopetrosis with renal tubular acidosis	99916	Ovarian malignant Sertoli-Leydig cell tumor	64686	Painful ophthalmoplegia
178389	Osteopetrosis-hypogammaglobulinemia syndrome	398987	Ovarian malignant teratoma	300501	Painful orbital and systemic neurofibromas-marfanoid habitus syndrome
94063	Osteopikilosis-short stature-intellectual disability syndrome	398961	Ovarian mucinous adenocarcinoma	90797	PAIS
2787	Osteoporosis-macrocephaly-blindness-joint hyperlaxity syndrome	99916	Ovarian Sertoli-Leydig cell cancer	477993	Palatal anomalies-multiple diastemata-facial dysmorphism-developmental delay syndrome
2786	Osteoporosis-oculocutaneous hypopigmentation syndrome	206473	Ovarian tumor of low malignant potential	477993	Palatal anomalies-widely spaced teeth-facial dysmorphism-developmental delay syndrome
2788	Osteoporosis-pseudoglioma syndrome	99853	Ovarioleukodystrophy	1388	Palatodigital syndrome, Cotel-Manzke type
666	Osteopsathyrosis	137634	Overgrowth-macrocephaly-facial dysmorphism syndrome	171695	Pallidopyramidal syndrome
668	Osteosarcoma	3203	Overhydrated hereditary stomatocytosis	3138	Pallister ulnar-mammary syndrome
2760	Osteosarcoma-limb anomalies-erythroid macrocytosis syndrome	206572	Overlap myositis	672	Pallister-Hall syndrome
178377	Osteosclerosis-developmental delay-craniosynostosis syndrome	326	Owren disease	884	Pallister-Killian syndrome
75325	Osteosclerosis-ichthyosis-premature ovarian failure syndrome	832	OXCT1 deficiency	2804	Pallister-W syndrome
2905	Osteosclerotic myeloma	31	Oxoglutaricaciduria	737	Palmar, plantar and disseminated porokeratosis
1338	Ostravik-Lindemann-Solberg syndrome	33572	Oxoprolinuria due to oxoprolinase deficiency	2184	Palmer-Pagon syndrome
99965	O'Sullivan-McLeod syndrome	79302	Oxysterol 7-alpha-hydroxylase deficiency	659	Palmopantar and periorificial keratoderma
664	OTC deficiency	36355	P2Y12 defect	34217	Palmopantar hyperkeratosis with arrhythmogenic cardiomyopathy
1308	OTCS	35664	P5CS deficiency	140966	Palmopantar hyperkeratosis, Nagashima type
2791	Otodental dysplasia	35120	P5N deficiency	50944	Palmopantar hyperkeratosis-cystic eyelids-hypodontia-hypotrichosis syndrome
2791	Otodental syndrome	98971	PACD	2202	Palmopantar hyperkeratosis-deafness syndrome
2792	Otofaciocervical syndrome	2796	Pachydermoperiostosis	2198	Palmopantar hyperkeratosis-esophageal carcinoma syndrome
141136	Otomandibular dysostosis	→2995	Pachygyria-epilepsy-intellectual disability-dysmorphism syndrome	2202	Palmopantar hyperkeratosis-hearing loss syndrome
141136	Otomandibular syndrome	2798	Pachygyria-intellectual disability-epilepsy syndrome	2342	Palmopantar hyperkeratosis-periodontopathia-onychogryposis syndrome
2793	Otoonychoperoneal syndrome	2309	Pachyonychia congenita	384	Palmopantar hyperkeratosis-sclerodactyly syndrome
669	Otopalatodigital syndrome	1952	Pacman dysplasia	2201	Palmopantar hyperkeratosis-spastic paralysis syndrome
90650	Otopalatodigital syndrome type 1	140989	PACNS	85112	Palmopantar hyperkeratosis-XX sex reversal-predisposition to squamous cell carcinoma syndrome
90652	Otopalatodigital syndrome type 2	706	PAD	1010	Palmopantar keratoderma and congenital alopecia, Stevanovic type
1427	Otospondylomegaepiphyseal dysplasia	477749	PADMAL	1366	Palmopantar keratoderma and congenital alopecia, Wallis type
457252	OTSCC	441	PAF	34217	Palmopantar keratoderma with arrhythmogenic cardiomyopathy
69082	OTUDP syndrome	95232	PAFAH1B1-related lissencephaly	→2199	Palmopantar keratoderma with tonotubular keratin
50943	Oudtshoorn disease	180275	Paget disease of the breast		
1179	Ouvrier-Billson syndrome	180275	Paget disease of the nipple		
213504	Ovarian adenocarcinoma	357131	Paget-Schrotter disease		
213512	Ovarian carcinosarcoma	52430	Pagetoid amyotrophic lateral sclerosis		
398971	Ovarian clear cell adenocarcinoma	52430	Pagetoid neuroskeletal syndrome		
314473	Ovarian fibroma	178517	Pagetoid reticulosis, Woringer-Kolopp type		
314478	Ovarian fibrothecoma	180275	Paget's disease of the nipple		
64739	Ovarian hyperstimulation syndrome	991	PAGOD syndrome		
398987	Ovarian immature teratoma	2802	Pagon-Bird-Detter syndrome		
		716	PAH deficiency		
		1993	Pai syndrome		
		37202	Painful bladder syndrome		
		324636	Painful bruising syndrome		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
140966	Palmoplantar keratoderma, Nagashima type	424080	Pancreatic undifferentiated carcinoma with osteoclast-like giant cells	684	Paramyotonia congenita of Von Eulenburg
86919	Palmoplantar keratoderma-clinodactyly syndrome		677	Pancreatoblastoma	2812
50944	Palmoplantar keratoderma-cystic eyelids-hypodontia-hypotrichosis syndrome	317473	Pancytopenia due to IKZF1 mutations	99889	Paraneoplastic Cushing syndrome
2202	Palmoplantar keratoderma-deafness syndrome	401764	Pancytopenia-developmental delay syndrome	1183	Paraneoplastic opsoclonus-myoclonus
2198	Palmoplantar keratoderma-esophageal carcinoma syndrome	66624	PANDAS	1183	Paraneoplastic opsoclonus-myoclonus-ataxia syndrome
2202	Palmoplantar keratoderma-hearing loss syndrome	95513	Panhypophysitis	63455	Paraneoplastic pemphigus
2342	Palmoplantar keratoderma-periodontopathia-onychogryposis syndrome	90695	Panhypopituitarism	71505	Paraneoplastic retinopathy
384	Palmoplantar keratoderma-sclerodactyly syndrome	97336	Panner disease	279928	Paraneoplastic uveitis
2201	Palmoplantar keratoderma-spastic paralysis syndrome	90159	Panniculitis and localized lipodystrophy	231445	Paraparetic variant of GBS
85112	Palmoplantar keratoderma-XX sex reversal-predisposition to squamous cell carcinoma syndrome	157850	Pantothenate kinase-associated neurodegeneration	231445	Paraparetic variant of Guillain-Barré syndrome
→79502	Palmoplantar porokeratosis of Mantoux	440427	PAP, Reunion island type	2823	Paraplegia-brachydactyly-cone-shaped epiphysis syndrome
163927	Palmoplantar pustulosis	69126	PAPA syndrome	2824	Paraplegia-intellectual disability-hyperkeratosis syndrome
767	PAN	213817	Papillary carcinoma of the cervix uteri	31827	Paraquat poisoning
98815	Panayiotopoulos syndrome	213726	Papillary carcinoma of the corpus uteri	2646	Parastremmatic dwarfism
424046	Pancreatic acinar cell carcinoma	208600	Papillary fibroelastoma of the heart	363478	Paratesticular adenocarcinoma
93292	Pancreatic adenoma	251962	Papillary glioneuronal tumor	143	Parathyroid carcinoma
65288	Pancreatic and cerebellar agenesis	146	Papillary or follicular thyroid carcinoma	443227	Paratyphoid fever
97282	Pancreatic cholera	319298	Papillary renal cell adenocarcinoma	2825	PARC syndrome
309108	Pancreatic colipase deficiency	319298	Papillary renal cell carcinoma	268826	Parietal encephalocele
2255	Pancreatic hypoplasia-diabetes-congenital heart disease syndrome	251915	Papillary tumor of the pineal region	251290	Parietal foramina with cleidocranial dysostosis
811	Pancreatic insufficiency and bone marrow dysfunction	1475	Papillo-renal syndrome	851	Parietal foramina with cleidocranial dysplasia
199337	Pancreatic insufficiency-anemia-hyperostosis syndrome	2807	Papilloma of choroid plexus	306674	Paris-Trousseau thrombocytopenia
424058	Pancreatic intraductal papillary mucinous carcinoma	2750	Papillon-Léage-Psaume syndrome	199351	PARK9
424053	Pancreatic mucinous cystadenocarcinoma	678	Papillon-Lefèvre syndrome	90307	PARK14
424080	Pancreatic osteoclastic giant cell tumor	86819	Papular atrichia	90307	Parkes Weber syndrome
97278	Pancreatic polypeptidoma	228264	Papular elastorrhexis	171695	Parkinsonian-pyramidal syndrome
424073	Pancreatic serous cystadenocarcinoma	313936	Papular epidermal nevi with skyline basal cell layers syndrome	314632	Parkinsonism due to ATP13A2 deficiency
424065	Pancreatic solid pseudopapillary carcinoma	90395	Papular mucinosis of infancy	178509	Parkinsonism with alveolar hypoventilation and mental depression
424039	Pancreatic squamous cell carcinoma	158008	Papular xanthoma	97355	Parkinsonism with dementia of Guadeloupe
309031	Pancreatic triacylglycerol lipase deficiency	679	Papulosis atrophican maligna	90020	Parkinsonism-dementia-ALS complex
309031	Pancreatic triglyceride lipase deficiency	464458	Paracetamol poisoning	90035	Paroxysmal cold hemoglobinuria
		99056	Parachute tricuspid valve	53583	Paroxysmal dystonic choreathetosis with episodic ataxia and spasticity
		73260	Paracoccidioidomycosis	98811	Paroxysmal exertion-induced dyskinesia
		97286	Paraganglioma and gastric stromal sarcoma	46348	Paroxysmal extreme pain disorder
		324299	Paraganglioma-somatostatinoma-polycythemia syndrome	157835	Paroxysmal hemicrania
		326	Parahemophilia	→98784	Paroxysmal hypnagogic dyskinesia
		306530	Parálisis facial hereditaria congénita con pérdida de audición variable	→98784	Paroxysmal hypnagogic dystonia
		306530	Parálisis facial hereditaria congénita con sordera variable	→98784	Paroxysmal hypnogenic dyskinesia
		141242	Paramedian nasal cleft	98809	Paroxysmal kinesigenic choreathetosis
		684	Paramyotonia congenita		

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98809	Paroxysmal kinesigenic dyskinesia	458785	Partially involuting congenital hemangioma	96192	Paternal uniparental disomy of chromosome 7
31709	Paroxysmal kinesigenic dyskinesia and infantile convulsions	85453	Partington disease	99324	Paternal uniparental disomy of chromosome 13
→98784	Paroxysmal nocturnal dyskinesia	94083	Partington syndrome	96334	Paternal uniparental disomy of chromosome 14
447	Paroxysmal nocturnal hemoglobinuria	→193	Partington-Anderson syndrome	96194	Paternal uniparental disomy of chromosome 20
98810	Paroxysmal non-kinesigenic dyskinesia	94083	Partington-Mulley syndrome	96195	Paternal uniparental disomy of chromosome 21
98810	Paroxysmic non-kinesigenic choreoathetosis	295	Parvovirus antenatal infection	261524	Paternal uniparental disomy of chromosome X
1214	Parry-Romberg syndrome	1394	Pascual-Castroviejo syndrome type 1	96194	Paternal UPD20
574	Partial 21q monosomy	42775	Pascual-Castroviejo syndrome type 2	2439	Patterson-Stevenson syndrome
79087	Partial acquired lipodystrophy	289478	PASH syndrome	2439	Patterson-Stevenson-Fontaine syndrome
2805	Partial agenesis of the pancreas	1252	Pashayan syndrome	79136	PATX
381	Partial albinism-immunodeficiency syndrome	1252	Pashayan-Prozansky syndrome	93126	Pauci-immune glomerulonephritis
90797	Partial androgen insensitivity syndrome	2278	Passwell-Goodman-Siprkowski syndrome	97563	Pauci-immune glomerulonephritis with ANCA
90797	Partial androgen resistance syndrome	3378	Patau syndrome	97563	Pauci-immune glomerulonephritis with antineutrophil cytoplasmic antibody
1330	Partial atrioventricular canal	→1509	Patella aplasia-coxa vara-tarsal synostosis syndrome	97564	Pauci-immune glomerulonephritis without ANCA
1330	Partial atrioventricular canal defect	86789	Patella aplasia/hypoplasia	97564	Pauci-immune glomerulonephritis without antineutrophil cytoplasmic antibody
1646	Partial chromosome Y deletion	295041	Patella aplasia/hypoplasia, bilateral	85410	Pauciarticular chronic arthritis
401959	Partial corpus callosum agenesis-cerebellar vermis hypoplasia with posterior fossa cysts syndrome	295038	Patella aplasia/hypoplasia, unilateral	247839	Pauciarticular chronic arthritis with anti-nuclear antibodies
98950	Partial cryptophthalmia	706	Patent arterial duct	247846	Pauciarticular chronic arthritis without anti-nuclear antibodies
90076	Partial deep dermal and full thickness burns	706	Patent arterial duct-bicuspid aortic valve-hand anomalies syndrome	1330	PAVC
79312	Partial deficiency of methylmalonyl-CoA mutase	46627	Patent ductus arteriosus	2038	PAVM
261318	Partial duplication of chromosome 20p	228190	Patent ductus arteriosus with facial dysmorphism and abnormal fifth digits	186	PBC
261318	Partial duplication of the short arm of chromosome 20	99108	Patent ductus arteriosus-bicuspid aortic valve-hand anomalies syndrome	75373	PBCRA
101046	Partial epilepsy with auditory aura	431341	Patent foramen ovale	289666	PBL
101046	Partial epilepsy with auditory features	254531	Patent urachus	2309	PC
2704	Partial facial palsy with urinary abnormalities	254525	Paternal 14q32.2 hypomethylation syndrome	54247	PCA
744	Partial gigantism-nevi-hemihypertrophy-macrocephaly syndrome	261304	Paternal 14q32.2 microdeletion syndrome	88628	PCARP
254693	Partial hydatidiform mole	261304	Paternal 20q13.2-q13.3 microdeletion syndrome	231426	PCB variant of GBS
79292	Partial LCAT deficiency	254525	Paternal 20q13.2q13.3 microdeletion syndrome	231426	PCB variant of Guillain-Barré syndrome
343	Partial mevalonate kinase deficiency	261304	Paternal del(14)(q32.2)	1578	PCBD deficiency
254693	Partial molar pregnancy	261304	Paternal del(20)(q13.2q13.3)	247198	PCCA
2805	Partial pancreatic agenesis	261304	Paternal monosomy 14q32.2	244	PCD
180129	Partial septate uterus	261304	Paternal monosomy 20q13.2-q13.3	178544	PCDLBCL,LT
157769	Partial situs inversus	261304	Paternal monosomy 20q13.2q13.3	178540	PCFCL
261318	Partial trisomy of chromosome 20p	251004	Paternal uniparental disomy of chromosome 1	90035	PCH
261318	Partial trisomy of the short arm of chromosome 20	96190	Paternal uniparental disomy of chromosome 5	2254	PCH1
		96191	Paternal uniparental disomy of chromosome 6	2524	PCH2
				97249	PCH3
				166063	PCH4
				166068	PCH5

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166073	PCH6	2836	PEHO syndrome	2880	PEPCK deficiency
284339	PCH7	99807	PEHO-like syndrome	2576	Perheentupa syndrome
324569	PCH8	48686	PEL	767	Periarteritis nodosa
369920	PCH9	702	Pelizaeus-Merzbacher brain sclerosis	2847	Pericardial and diaphragmatic defect
97249	PCH with optic atrophy			2576	Pericardial constriction-growth failure syndrome
97249	PCH without dyskinesia	280229	Pelizaeus-Merzbacher disease in female carriers		
411493	PCH10			280210	Pelizaeus-Merzbacher disease type II
71528	PCI deficiency	280219	Pelizaeus-Merzbacher disease, classic form		
454714	PCL			280210	Pelizaeus-Merzbacher disease, connatal form
2924	PCLD	280234	Pelizaeus-Merzbacher disease, null syndrome		
178536	PCMZL			280224	Pelizaeus-Merzbacher disease, transitional form
438134	PCNA-related progressive neurodegenerative photosensitivity syndrome	280270	Pelizaeus-Merzbacher-like disease		
46135	PCNSL			280293	Pelizaeus-Merzbacher-like disease due to AIMP1 mutation
140989	PCNSV	280282	Pelizaeus-Merzbacher-like disease due to GJC2 mutation		
101330	PCT			280288	Pelizaeus-Merzbacher-like disease due to HSPD1 mutation
163746	PCWH	97352	Pellagra		
90020	PDALS	2837	Pellagra-like skin rash-neurological manifestations syndrome	342	Periodic disease
293462	PDCD	137672	Pellucid marginal degeneration	42642	Periodic fever-aphtous stomatitis-pharyngitis-adenopathy syndrome
289157	PDDR1	2840	Pelvic dysplasia-arthrogryposis of lower limbs syndrome	436166	Periodic fever-infantile enterocolitis-autoinflammatory syndrome
439822	PDE4D haploinsufficiency syndrome				
765	PDH	2839	Pelvis-shoulder dysplasia	397750	Periodic paralysis with later-onset distal motor neuropathy
79246	PDH phosphatase deficiency	93333	Pelviscapular dysplasia		
79243	PDHAD	63275	Pemphigoid gestationis	79136	Periodic vestibulocerebellar ataxia
255138	PDHBD	79480	Pemphigus erythematosus		
765	PDHC	79481	Pemphigus foliaceus	563	Peripartum cardiomyopathy
2796	PDP	79479	Pemphigus vegetans		
85453	PDR	704	Pemphigus vulgaris	1795	Peripheral dysostosis
75496	PDS	994	Pena-Shokeir syndrome type 1		
699	Pearson syndrome	1466	Pena-Shokeir syndrome type 2	2400	Peripheral motor neuropathy-dysautonomia syndrome
2835	Pectus excavatum-macrocephaly-dysplastic nails syndrome	705	Pendred syndrome		
98811	PED	398053	Penile adenocarcinoma	213812	Peripheral neuroectodermal cancer of cervix uteri
439175	Pediatric AIS	49	Penile agenesis		
439175	Pediatric arterial ischemic stroke	398058	Penile squamous cell carcinoma	90120	Peripheral neuropathy and optic atrophy
66624	Pediatric autoimmune disorders associated with Streptococcus infections	49	Penosclerotic transposition		
		93682	Pediatric Castleman disease	11	Penta-X
33402	Pediatric HCC	2842	Penosclerotic transposition	1335	Pentalogy of Cantrell
33402	Pediatric hepatocellular carcinoma	313936	PENS syndrome	11	Pentasomy X
477738	Pediatric multiple sclerosis	11	Penta-X	2843	Pentosuria
93552	Pediatric systemic lupus erythematosus	352447	PEO-myopathy-emaciation syndrome	2905	PEP syndrome
263548	Peeling skin syndrome type A	2905	PEP syndrome		
263553	Peeling skin syndrome type B	2843	Pentosuria		
444138	Peeling skin-leukonuchia-acral punctate keratoses-cheilitis-knuckle pads syndrome	352447	PEO-myopathy-emaciation syndrome		
		2905	PEP syndrome		

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171848	Peripheral neuropathy, Fiskerstrand type	708	Peters anomaly	2874	Phakomatosis pigmentokeratotica
397744	Peripheral neuropathy-myopathy-hoarseness-deafness syndrome	709	Peters anomaly with short limb dwarfism	2875	Phakomatosis pigmentovasularis
		101033	Peters anomaly-cataract syndrome	79483	Phakomatosis pigmentovasularis type 2
397744	Peripheral neuropathy-myopathy-hoarseness-hearing loss syndrome	708	Peters congenital glaucoma	79485	Phakomatosis pigmentovasularis type 3
		709	Peters plus syndrome		
370348	Peripheral PNET	2776	Petit-Fryns syndrome	79484	Phakomatosis pigmentovasularis type 5
370348	Peripheral primitive neuroectodermal tumor	2963	Petty syndrome	79485	Phakomatosis spilorosea
		2963	Petty-Laxova-Wiedemann syndrome		
97927	Peripheral resistance to thyroid hormones	2869	Peutz-Jeghers syndrome	352636	Phalangeal microgeodic syndrome
		42642	PFAPA syndrome	352636	Phalangeal osteolysis
168816	Peritoneal cystic mesothelioma	1980	PFBC	171848	PHARC syndrome
171676	Periventricular leukomalacia	90042	PFCP	231426	Pharyngeal-cervical-brachial variant of Guillain-Barré syndrome
98892	Periventricular nodular heterotopia	412206	PFE	231426	Pharyngeal-cervical-brachial weakness
2849	Perlman syndrome	710	Pfeiffer syndrome		
438266	PERM	93258	Pfeiffer syndrome type 1	231426	Pharyngo-cervico-brachial variant of GBS
99885	Permanent neonatal diabetes mellitus	93259	Pfeiffer syndrome type 2		
		93260	Pfeiffer syndrome type 3	231426	Pharyngo-cervico-brachial variant of Guillain-Barré syndrome
65288	Permanent neonatal diabetes mellitus-pancreatic and cerebellar agenesis syndrome	3224	Pfeiffer-Kapferer syndrome		
		2921	Pfeiffer-Mayer syndrome		
2850	Perniola-Krajewska-Carnevale syndrome	2871	Pfeiffer-Palm-Teller syndrome	228410	PHD syndrome
		2872	Pfeiffer-Singer-Zschesche syndrome	48652	Phelan-McDermid syndrome
2971	Peroxisomal acyl-CoA oxidase deficiency	33577	Pfeiffer-Weber-Christian syndrome	1919	Phenobarbital embryopathy
		2019	PFFD	84064	Phenotypic diarrhea
93598	Peroxisomal alanine-glyoxylate aminotransferase deficiency	172	PFIC	716	Phenylalanine hydroxylase deficiency
		79306	PFIC1		
2855	Perrault syndrome	79304	PFIC2	716	Phenylketonuria
75374	PERRS	79305	PFIC3	226	Phenylketonuria type 2
178509	Perry syndrome	480483	PFIC4	2209	Phenylketonuric embryopathy
99120	Persistent eustachian valve	480476	PFIC5	1912	Phenytoin embryofetopathy
91495	Persistent fetal vasculature syndrome	91495	PFVS	→168569	PHID
		397937	PGBM1	75508	Phlebotactic osteohypoplastic angiodysplasia
99076	Persistent fifth aortic arch	319646	PGM1-CDG		
91495	Persistent hyperplastic primary vitreous	443811	PGM3-CDG	294975	Phocomelia
		443811	PGM3-related congenital disorder of glycosylation		
398147	Persistent idiopathic facial pain	251962	PGNT	2879	Phocomelia, Schinzel type
99109	Persistent left superior caval vein connecting to the left-sided atrium	1214	PHA	2878	Phocomelia-ectrodactyly-deafness-sinus arrhythmia syndrome
		757	PHA2		
99109	Persistent left superior vena cava connecting to the left-sided atrium	88938	PHA2A	3439	Phocomelia-thrombocytopenia-encephalocele-urogenital malformations syndrome
		88939	PHA2B		
99109	Persistent left SVC connecting to the left-sided atrium	88940	PHA2C	534	Phosphatidylinositol 4,5-bisphosphate 5-phosphatase deficiency
		300525	PHA2D		
2856	Persistent Müllerian derivatives	300530	PHA2E	2880	Phosphoenolpyruvate carboxykinase deficiency
2856	Persistent Müllerian duct syndrome	756	PHA type 1		
706	Persistent patency of the arterial duct	42775	PHACE syndrome	436	Phosphoethanolaminuria
97341	Persistent placoid maculopathy	209959	Phacoallergic endophthalmitis	→319646	Phosphoglucomutase 1 deficiency
300324	Persistent polyclonal B-cell lymphocytosis	209959	Phacoanaphylactic uveitis	35069	Phospholipase A2-associated neurodegeneration
		209959	Phacoantigenic endophthalmitis		
300324	Persistent polyclonal B-cell lymphocytosis with binucleated lymphocytes	757	PHAI1	79318	Phosphomannomutase 2 deficiency
		209959	Phako-anaphylactic endophthalmitis	79319	Phosphomannose isomerase deficiency
2380	Perthes disease	79483	Phakomatosis cesioflammea		
1489	Pertussis	79484	Phakomatosis cesiomarmorata		

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3222	Phosphoribosylpyrophosphate synthetase superactivity	447961	Pigmentation defects-palmoplantar keratoderma-skin carcinoma syndrome	91351	Pituitary dermoid and epidermoid cysts
284417	Phosphoserine aminotransferase deficiency			99725	Pituitary gigantism
166409	Photosensitive epilepsy	→168569	Pigmented hypertrichosis with insulin-dependent diabetes mellitus syndrome	2965	Pituitary lactotrophic adenoma
91495	PHPV			95496	Pituitary stalk interruption syndrome
30924	PHSH	251295	Pigmented paravenous retinochoroidal atrophy	91347	Pituitary thyrotrophic adenoma
180261	Phylloide tumor			96253	Pituitary-dependent Cushing syndrome
180261	Phylloide tumor	66627	Pigmented villonodular synovitis	2897	Pityriasis rubra pilaris
773	Phytanic-CoA hydroxylase deficiency	280633	PIGN-CDG		
2882	Phytosterolemia	480506	PIHL	1078	Piussan-Lenaerts-Mathieu syndrome
→33364	PIBIDS syndrome	169	Pili annulati	2869	PJS
505	Piccardi-Lassueur-Little syndrome	720	Pili bifurcati	157850	PKAN
2885	Piebald trait-neurologic defects syndrome	79492	Pili gemini	216873	PKAN, atypical form
2884	Piebaldism	79492	Pili multigemini	216866	PKAN, classic form
→1263	Piepkorn dysplasia	2889	Pili torti	238455	PKDYS
1566	Pierquin syndrome	2891	Pili torti-developmental delay-neurological abnormalities syndrome	716	PKU
2886	Pierre Robin sequence-congenital heart defect-talipes syndrome			2890	Pili torti-onychodysplasia syndrome
2888	Pierre Robin sequence-faciodigital anomaly syndrome	1410	Pili trianguli et canaliculi	477787	PLA2G4A-related platelet dysfunction
3450	Pierre Robin sequence-fetal chondrodysplasia syndrome	2741	Pillay syndrome	199351	PLA2G6-related dystonia-parkinsonism
1388	Pierre Robin sequence-hyperphalangy-clinodactyly syndrome	251612	Pilocytic astrocytoma	439167	Placental insufficiency
		2892	Pilodental dysplasia-refractive errors syndrome	99928	Placental site trophoblastic tumor
		91414	Pilomatricoma	444138	PLACK syndrome
3104	Pierre Robin sequence-oligodactyly syndrome	228379	Pilomatrix dysplasia	707	Plague
		91414	Pilomatrixoma	300359	PLAID
2886	Pierre Robin syndrome-congenital heart defect-talipes syndrome	251615	Pilomyxoid astrocytoma	79141	Plamoplantar hyperkeratosis nummularis
2888	Pierre Robin syndrome-faciodigital anomaly syndrome	2894	Pilotto syndrome	79141	Plamoplantar keratoderma nummularis
		251919	Pineal parenchymal tumor of intermediate differentiation	35069	PLAN
3450	Pierre Robin syndrome-fetal chondrodysplasia syndrome	251909	Pineoblastoma	199251	Plantar fibromatosis
		251912	Pineocytoma	251515	Plantar flexion contracture
1388	Pierre Robin syndrome-hyperphalangy-clinodactyly syndrome	49382	Pingelapese blindness	158769	Plaque-form urticaria pigmentosa
		3353	Pinheiro-Freire Maia-Miranda syndrome	454714	Plasma cell leukemia
		247165	Pink disease	29073	Plasma cell myeloma
2670	Pierson syndrome	155838	Pinnae fistula or cyst	329	Plasma thromboplastin antecedent deficiency
398147	PIFP	→2510	Pinsky-Di George-Harley syndrome		
217557	PIG	279904	PIOL	289666	Plasmablastic lymphoma
99908	Pigeon-breeder lung disease	→79189	Pipecolic acidemia	86855	Plasmacytoma
3474	PIGL-CDG	2896	Pitt-Hopkins syndrome	722	Plasminogen deficiency type 1
83639	PIGM-CDG	221150	Pitt-Hopkins-like syndrome	439881	Plastic bronchitis
978	Pigment anomaly-ectrodactyly-hypodontia syndrome	→280	Pitt-Rogers-Danks syndrome	721	Platelet alpha-granule deficiency
		93395	Pitt-Williams brachydactyly	477787	Platelet dysfunction due to cytosolic phospholipase-A2 alpha deficiency
999	Pigmentary disorder with hearing loss	251623	Pituicytoma	52530	Platelet type-von Willebrand disease
64755	Pigmentary hairy epidermal nevus	95613	Pituitary apoplexy	79434	Platinum oculocutaneous albinism
435	Pigmentary mosaicism, Ito type	300385	Pituitary carcinoma		
313808	Pigmentary orthochromatic leukodystrophy	96253	Pituitary corticotroph micro-adenoma	85166	Platyspondylic dysplasia, Torrance type
		91354	Pituitary deficiency due to empty sella turcica syndrome	85166	Platyspondylic dysplasia, Torrance-Luton type
→193	Pigmentary retinopathy-intellectual disability syndrome	91350	Pituitary deficiency due to Rathke's pouch cysts		

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85166	Platyspondylic lethal skeletal dysplasia, Torrance type	79318	PMM2-CDG	247854	Polyarthritis without rheumatoid factor with anti-nuclear antibodies
		26790	PMP		
2899	Platyspondyly-amelogenesis imperfecta syndrome	476394	PMP2-related Charcot-Marie-Tooth disease type 1	247861	Polyarthritis without rheumatoid factor without anti-nuclear antibodies
300359	PLCG2-associated antibody deficiency and immune dysregulation	476394	PMP2-related Charcot-Marie-Tooth neuropathy type 1	450322	Polyclonal hyperviscosity syndrome
		476394	PMP2-related CMT1		
137810	PLCNA	476394	PMP2-related hereditary motor and sensory neuropathy type 1	2770	Polycystic lipomembranous osteodysplasia with sclerosing leukoencephalopathy
99969	Pleomorphic liposarcoma	477817	PMP22-RAI1 contiguous gene duplication syndrome	2795	Polycystic ovaries-urethral sphincter dysfunction syndrome
293199	Pleomorphic rhabdomyosarcoma	99885	PNDM	729	Polycythemia rubra vera
454821	Pleomorphic salivary gland adenoma	64741	Pneumoblastoma	729	Polycythemia vera
251607	Pleomorphic xanthoastrocytoma	55655	Pneumococcal meningitis	93339	Polydactyly of a biphalangal thumb
449266	Pleural empyema	723	Pneumocystosis	295146	Polydactyly of a biphalangal thumb, bilateral
50251	Pleural mesothelioma	90066	Pneumonia caused by Pseudomonas aeruginosa infection	295144	Polydactyly of a biphalangal thumb, unilateral
99131	Pleuro-pericardial cyst	447	PNH	93336	Polydactyly of a triphalangal thumb
284343	Pleuro-pulmonary blastoma familial tumor susceptibility syndrome	760	PNP deficiency	295150	Polydactyly of a triphalangal thumb, bilateral
64742	Pleuropulmonary blastoma	760	PNPase deficiency	295148	Polydactyly of a triphalangal thumb, unilateral
284343	Pleuropulmonary blastoma familial tumor susceptibility syndrome	79096	PNPO deficiency	93337	Polydactyly of an index finger
99933	Pleuropulmonary blastoma type 1	79096	PNPO-related neonatal epileptic encephalopathy	295154	Polydactyly of an index finger, bilateral
99934	Pleuropulmonary blastoma type 2	246	POADS	295152	Polydactyly of an index finger, unilateral
99935	Pleuropulmonary blastoma type 3	2905	POEMS syndrome	2919	Polydactyly postaxial with median cleft of upper lip
280356	PLIN1-related familial partial lipodystrophy	2762	POH	2754	Polydactyly-cleft lip/palate-psycho motor retardation syndrome
280356	PLIN1-related FPLD	2908	Poikiloderma of Kindler	2917	Polydactyly-myopia syndrome
2770	PLO-SL	2909	Poikiloderma of Rothmund-Thomson	180229	Polyembryoma
2770	PLOSL	221008	Poikiloderma of Rothmund-Thomson type 1	453533	Polyendocrine-polyneuropathy syndrome
2375	Plott syndrome	221016	Poikiloderma of Rothmund-Thomson type 2	93308	Polyepiphyseal dysplasia type 1
280234	PLP1 null syndrome	221046	Poikiloderma with neutropenia	93307	Polyepiphyseal dysplasia type 4
678	PLS	221046	Poikiloderma with neutropenia, Clericuzio type	93311	Polyepiphyseal dysplasia type 5
35689	PLS	221046	Poikiloderma with neutropenia, Clericuzio type	397937	Polyglucosan body myopathy type 1
99969	PLS	2825	Poikiloderma-alopecia-retrognathism-cleft palate syndrome	456369	Polyglucosan body myopathy type 2
85166	PLSD-T	279947	POIS	180182	Polymastia
330015	Plumbism	130	Pokkuri death syndrome	447877	Polymerase proofreading-related adenomatous polyposis
54028	Plummer-Vinson syndrome	2911	Poland anomaly	300573	Polymicrogyria due to TUBB2B mutation
732	PM	2911	Poland sequence	250972	Polymicrogyria with optic nerve hypoplasia
764	PM	2911	Poland syndrome	2925	Polymicrogyria-turricephaly-hypogenitalism syndrome
454706	PMA	313808	POLD	64745	Polymorphic eruption of pregnancy
702	PMD	2912	Poliomyelitis	1243	Polymorphic vitelline macular degeneration
2856	PMDS	330009	Poliomyelitis in patients with immunodeficiencies deemed at risk	93569	Polymyalgia rheumatica
308	PME type 1	→33364	Pollitt syndrome		
501	PME type 2	11	Poly-X		
263516	PME type 3	767	Polyarteritis nodosa		
402082	PME type 5	29207	Polyarthritis enterica		
280620	PME type 6	85435	Polyarthritis with rheumatoid factor		
435438	PME type 7	85408	Polyarthritis without rheumatoid factor		
424027	PME type 8				
457265	PME type 9				
352596	PMED				
217260	PML				
280270	PMLD				
280282	PMLD1				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
732	Polymyositis			295008	Postaxial polydactyly of foot
639	Polyneuropathy associated with IgM monoclonal gammopathy with anti-MAG	284339	Pontocerebellar hypoplasia-46,XY disorder of sex development syndrome	295008	Postaxial polydactyly of toes
2905	Polyneuropathy-endocrinopathy-plasma cell dyscrasia syndrome	284400	Poorly differentiated neuroendocrine carcinoma of the bladder	295181	Postaxial polydactyly of toes, bilateral
2926	Polyneuropathy-hand defect syndrome	213777	Poorly differentiated neuroendocrine carcinoma of the cervix uteri	295179	Postaxial polydactyly of toes, unilateral
171848	Polyneuropathy-hearing loss-ataxia-retinitis pigmentosa-cataract syndrome	213777	Poorly differentiated neuroendocrine carcinoma of the corpus uteri	93334	Postaxial polydactyly type A
2928	Polyneuropathy-intellectual disability-acromicria-premature menopause syndrome	213731	Poorly differentiated neuroendocrine carcinoma of the endometrium	295165	Postaxial polydactyly type A, bilateral
93276	Polyostotic fibrous dysplasia	213777	Poorly differentiated neuroendocrine cervical carcinoma	295163	Postaxial polydactyly type A, unilateral
160148	Polypoid prolapsing folds	263339	Poorly differentiated thymic neuroendocrine carcinoma	93335	Postaxial polydactyly type B
2869	Polyps and spots syndrome	1300	Popliteal web syndrome	295169	Postaxial polydactyly type B, bilateral
208981	Polyradiculoneuropathy associated with IgG/IgA/IgM monoclonal gammopathy without known antibodies	95699	POR deficiency	295167	Postaxial polydactyly type B, unilateral
141091	Polyrhinia	666	Porak and Durante disease	420584	Postaxial polydactyly-anterior pituitary anomalies-facial dysmorphism syndrome
141091	Polyrrhinia	95699	PORD	2916	Postaxial polydactyly-dental and vertebral anomalies syndrome
93338	Polysyndactyly	2940	Porencephaly	2920	Postaxial polydactyly-intellectual disability syndrome
295161	Polysyndactyly, bilateral	2941	Porencephaly-cerebellar hypoplasia-internal malformations syndrome	93406	Postaxial syndactyly with metacarpal synostosis
93405	Polysyndactyly, Haas type	306547	Porencephaly-microcephaly-bilateral congenital cataract syndrome	2730	Postaxial tetramelic oligodactyly
295159	Polysyndactyly, unilateral	370022	Poretti-Boltshauser syndrome	263352	Postcardiotomy right ventricular failure
2934	Polysyndactyly-cardiac malformation syndrome	735	Porokeratosis of Mibelli	97349	Postencephalitic parkinsonism
228410	Polyvalvular heart disease syndrome	737	Porokeratosis plantaris palmaris et disseminata	98971	Posterior amorphous corneal dystrophy
139426	POMA	166286	Porokeratotic eccrine nevus	98971	Posterior amorphous stromal dystrophy
1183	POMA syndrome	166286	Porokeratotic eccrine ostial and dermal duct nevus	88628	Posterior column ataxia-retinitis pigmentosa syndrome
71526	POMC deficiency	101330	Porphyria cutanea tarda	54247	Posterior cortical atrophy
365	Pompe disease	443057	Porphyria cutanea tarda type I	2064	Posterior fusion of lumbosacral vertebrae-blepharoptosis syndrome
308552	Pompe disease, infantile onset	443062	Porphyria cutanea tarda type II	95706	Posterior hypospadias
420429	Pompe disease, late onset	100924	Porphyria due to ALA dehydratase deficiency	268810	Posterior meningocele
99748	Pontiac fever	100924	Porphyria due to ALAD deficiency	98973	Posterior polymorphous corneal dystrophy
477749	Pontine autosomal dominant microangiopathy with leukoencephalopathy	100924	Porphyria due to delta-aminolevulinate dehydratase deficiency	98973	Posterior polymorphous dystrophy
269229	Pontine tegmental cap dysplasia	100924	Porphyria of Doss	93110	Posterior urethral valve
324569	Pontocerebellar hypoplasia due to CHMP1A mutation	79473	Porphyria variegata	48435	Postinfectious vasculitis
2254	Pontocerebellar hypoplasia type 1	100924	Port-wine nevi-mega cisterna magna-hydrocephalus syndrome	216452	Postlingual non-syndromic genetic deafness
2524	Pontocerebellar hypoplasia type 2	2703	Post-transplant lymphoproliferative disease	477673	Postnatal microcephaly-infantile hypotonia-spastic diplegia-dysarthria-intellectual disability syndrome
97249	Pontocerebellar hypoplasia type 3	70568	Postganglionic sepsis secondary to oropharyngeal infection	279947	Postorgasmic illness syndrome
166063	Pontocerebellar hypoplasia type 4	137839	Postaxial acrodysostosis	563	Postpartum cardiomyopathy
166068	Pontocerebellar hypoplasia type 5	246	Postaxial acrofacial dysostosis	443173	Postpartum psychosis
166073	Pontocerebellar hypoplasia type 6	246	Postaxial acrofacial dysostosis	2942	Postpolio sequelae
284339	Pontocerebellar hypoplasia type 7				
324569	Pontocerebellar hypoplasia type 8				
369920	Pontocerebellar hypoplasia type 9				
411493	Pontocerebellar hypoplasia type 10				

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2942	Postpolio syndrome	370348	PPNET	295152	Preaxial polydactyly type 3, unilateral
2942	Postpoliomyelitic syndrome	97278	PPoma	93338	Preaxial polydactyly type 4
2942	Postpoliomyelitis sequelae	163927	PPP	295161	Preaxial polydactyly type 4, bilateral
2942	Postpoliomyelitis syndrome	308013	PPP3 without elastoidosis	295159	Preaxial polydactyly type 4, unilateral
98913	Postsynaptic congenital myasthenic syndromes	79502	PPPP	2921	Preaxial polydactyly-colobomata-intellectual disability syndrome
		251295	PPRCA	1309	Precalcical canalicular ectasia
163921	Posttransplant acute limbic encephalitis	398980	PPSPC	99860	Precursor B-cell acute lymphoblastic leukemia
		324977	PRAAS	99860	Precursor B-cell acute lymphoblastic leukemia/lymphoma
443236	Postural tachycardia syndrome due to NET deficiency	739	Prader-Labhart-Willi syndrome	99860	Precursor B-cell acute lymphocytic leukemia
238606	POT	3409	Prader-Willi habitus-osteopenia-camptodactyly syndrome	99860	Precursor B-cell acute lymphocytic leukemia
→682	Potassium-sensitive normokalemic periodic paralysis	739	Prader-Willi syndrome	99860	Precursor B-cell acute lymphocytic leukemia/lymphoma
640	Potato-grubbing palsy	177910	Prader-Willi syndrome due to imprinting mutation	99860	Precursor B-cell acute lymphocytic leukemia
1713	Potocki-Lupski syndrome		Prader-Willi syndrome due to maternal uniparental disomy of chromosome 15	99860	Precursor B-cell acute lymphocytic leukemia/lymphoma
52022	Potocki-Shaffer syndrome	98754	Prader-Willi syndrome due to maternal uniparental disomy of chromosome 15	99861	Precursor T-cell acute lymphoblastic leukemia
3316	Potter sequence-cleft lip/palate-cardiopathy syndrome	98793	Prader-Willi syndrome due to paternal 15q11q13 deletion	99861	Precursor T-cell acute lymphoblastic leukemia/lymphoma
217067	Pouchitis		Prader-Willi syndrome due to paternal deletion of 15q11q13 type 1	99861	Precursor T-cell acute lymphocytic leukemia
2876	Powell-Chandra-Saal syndrome	177901	Prader-Willi syndrome due to paternal deletion of 15q11q13 type 1	99861	Precursor T-cell acute lymphocytic leukemia/lymphoma
2201	Powell-Venencie-Gordon syndrome		Prader-Willi syndrome due to paternal deletion of 15q11q13 type 2	99861	Precursor T-cell acute lymphocytic leukemia/lymphoma
314566	PPAOS	177904	Prader-Willi syndrome due to paternal deletion of 15q11q13 type 2	457088	Predisposition to invasive fungal disease due to CARD9 deficiency
447877	PPAP	398069	Prader-Willi syndrome due to point mutation	275555	Preeclampsia
79083	PPARG-related familial partial lipodystrophy		Prader-Willi syndrome due to translocation	69665	Pregnancy-related cholestasis
79083	PPARG-related FPLD	177907	Prader-Willi syndrome due to translocation	216445	Prelingual non-syndromic genetic deafness
284343	PPB familial tumor susceptibility syndrome	398073	Prader-Willi-like syndrome	276432	Premature aging appearance-developmental delay-cardiac arrhythmia syndrome
284343	PPBFTDS	171829	Prader-Willi-like syndrome due to deletion 6q16	363665	Premature aging syndrome, Penttinen type
300324	PPBL	398079	Prader-Willi-like syndrome due to point mutation	52183	Premature chromosome condensation with microcephaly and intellectual disability
168829	PPC	2956	Prata-Liberal-Goncalves syndrome	95486	Premature closure of the arterial duct
98973	PPCD	293462	Pre-Descemet corneal dystrophy	95486	Premature closure of the patent ductus arteriosus
93339	PPD1	245	Preaxial acrodysostosis	2114	Premature degenerative osteoarthropathy of the hip
93336	PPD2	2957	Preaxial deficiency-postaxial polydactyly-hypospadias syndrome	247638	Prenatal benign hypophosphatasia
93337	PPD3	295006	Preaxial polydactyly of foot	247638	Prenatal benign phosphoethanolaminuria
93338	PPD4	295006	Preaxial polydactyly of toes	247638	Prenatal benign Rathburn disease
75567	PPFG	295177	Preaxial polydactyly of toes, bilateral	90160	Pressure-induced localized lipomatrophy
411696	PPI-REE	295175	Preaxial polydactyly of toes, unilateral	98914	Presynaptic congenital myasthenic syndromes
411696	PPI-responsive esophageal eosinophilia	93339	Preaxial polydactyly type 1	79410	Pretibial DEB
411696	PPIRee	295146	Preaxial polydactyly type 1, bilateral	79410	Pretibial dystrophic epidermolysis bullosa
494	PPK mutilans and deafness	295144	Preaxial polydactyly type 1, unilateral		
79141	PPK nummularis	93336	Preaxial polydactyly type 2		
86923	PPK, Gamborg-Nielsen type	295150	Preaxial polydactyly type 2, bilateral		
140966	PPK, Nagashima type	295148	Preaxial polydactyly type 2, unilateral		
1010	PPK-CA, Stevanovic type	93337	Preaxial polydactyly type 3		
1366	PPK-CA, Wallis type	295154	Preaxial polydactyly type 3, bilateral		
2202	PPK-deafness syndrome				
79501	PPKP1				
79502	PPKP2				
38	PPKP3				
308013	PPKP3 without elastoidosis				
3077	PPM-X				
189439	PPNAD				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
2958	Prieto-Badia-Mulas syndrome	178536	Primary cutaneous marginal zone B-cell lymphoma	30924	Primary hypomagnesemia with secondary hypocalcemia
1451	Prieur-Griscelli syndrome			75391	Primary immunodeficiency due to MCM4 deficiency
930	Primary achalasia	86885	Primary cutaneous peripheral T-cell lymphoma NOS	90023	Primary immunodeficiency syndrome due to p14 deficiency
75564	Primary acquired sideroblastic anemia	86885	Primary cutaneous peripheral T-cell lymphoma not otherwise specified	90023	Primary immunodeficiency syndrome with short stature
85138	Primary Addison's disease	451602	Primary cutaneous plasmacytosis	447731	Primary immunodeficiency with multifaceted aberrant lymphoid immunity
874	Primary adult heart tumor	86885	Primary cutaneous unspecified peripheral T-cell lymphoma	75391	Primary immunodeficiency with natural-killer cell deficiency and adrenal insufficiency
85443	Primary amyloidosis	98807	Primary dystonia with mixed phenotype	431166	Primary immunodeficiency with post-measles-mumps-rubella vaccine viral infection
228272	Primary anetoderma	99657	Primary dystonia, DYT2 type	431166	Primary immunodeficiency with post-MMR vaccine viral infection
140989	Primary angiitis of the central nervous system	98805	Primary dystonia, DYT4 type	73272	Primary insulin-like growth factor deficiency
1572	Primary antibody deficiency	98806	Primary dystonia, DYT6 type	90362	Primary intestinal lymphangiectasia
2285	Primary basilar invagination	98807	Primary dystonia, DYT13 type	480506	Primary intrahepatic lithiasis
189427	Primary bilateral macronodular adrenal hyperplasia	370103	Primary dystonia, DYT17 type	458768	Primary intralymphatic angioendothelioma
186	Primary biliary cholangitis	306734	Primary dystonia, DYT21 type	279904	Primary intraocular lymphoma
186	Primary biliary cirrhosis	464440	Primary dystonia, DYT27 type	279904	Primary intraocular non-Hodgkin's lymphoma
779	Primary biliary cirrhosis and systemic scleroderma	48686	Primary effusion lymphoma	140436	Primary intraosseous venous malformation
314684	Primary bone lymphoma	90026	Primary erythralgia	137926	Primary laryngeal lymphangioma
46135	Primary brain lymphoma	357220	Primary essential cutis verticis gyrata	35689	Primary lateral sclerosis
300865	Primary C-ALCL	412206	Primary failure of tooth eruption	314709	Primary localized amyloidosis
267	Primary calpainopathy	98957	Primary familial amyloidosis of the cornea	137810	Primary localized cutaneous nodular amyloidosis
169464	Primary CD59 deficiency	90042	Primary familial and congenital polycythemia	319667	Primary lymphoid conjunctival tumor
46135	Primary central nervous system lymphoma	1980	Primary familial brain calcification	319667	Primary lymphoma of the conjunctiva
140989	Primary central nervous system vasculitis	90042	Primary familial polycythemia	228272	Primary macular atrophy
244	Primary ciliary dyskinesia	3337	Primary Fanconi renotubular syndrome	168811	Primary malignant peritoneal mesothelioma
→244	Primary ciliary dyskinesia, Kartagener type	3337	Primary Fanconi syndrome	98838	Primary mediastinal clear cell lymphoma of B-cell type
247522	Primary ciliary dyskinesia-retinitis pigmentosa syndrome	633	Primary GH insensitivity	98838	Primary mediastinal large B-cell lymphoma
46135	Primary CNS lymphoma	633	Primary GH resistance	238642	Primary megaureter, adult-onset form
477781	Primary condylar hyperplasia	633	Primary growth hormone insensitivity	252050	Primary melanoma of the central nervous system
90042	Primary congenital erythrocytosis	633	Primary growth hormone resistance	54370	Primary membranoproliferative glomerulonephritis
98976	Primary congenital glaucoma	100085	Primary hepatic neuroendocrine carcinoma	306558	Primary microcephaly-epilepsy-permanent neonatal diabetes syndrome
91138	Primary cryoglobulinemia	480506	Primary hepatolithiasis		
178528	Primary cutaneous aggressive epidermotropic CD8+ T-cell lymphoma	314950	Primary HES		
300865	Primary cutaneous anaplastic large cell lymphoma	314950	Primary hypereosinophilic syndrome		
178522	Primary cutaneous CD4+ small/medium-sized pleomorphic T-cell lymphoma	2232	Primary hypergonadotropic hypogonadism-partial alopecia syndrome		
178544	Primary cutaneous diffuse large B-cell lymphoma, leg type	682	Primary hyperkalemic periodic paralysis		
178528	Primary cutaneous epidermotropic cytotoxic CD8+ T-cell lymphoma	416	Primary hyperoxaluria		
178540	Primary cutaneous follicle center lymphoma	93598	Primary hyperoxaluria type 1		
178533	Primary cutaneous gamma/delta-positive T-cell lymphoma	93599	Primary hyperoxaluria type 2		
		93600	Primary hyperoxaluria type 3		
		682	Primary hyperPP		
		33208	Primary hypersomnia		
		1572	Primary hypogammaglobulinemia		

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391408	Primary microcephaly-mild intellectual disability-young-onset diabetes syndrome	263317	Primary thymic epithelial tumor type B	79087	Progressive cephalothoracic lipodystrophy
824	Primary myelofibrosis	98807	Primary torsion dystonia with predominant craniocervical or upper limb onset	247198	Progressive cerebello-cerebral atrophy
357225	Primary non-essential cutis verticis gyrata			1871	Progressive cone dystrophy
289356	Primary non-gestational choriocarcinoma of ovary	231580	Primary unilateral adrenal hyperplasia	220393	Progressive cutaneous systemic sclerosis
289356	Primary non-gestational ovarian choriocarcinoma	140989	Primary vasculitis of the central nervous system	220393	Progressive cutaneous systemic sclerosis
279897	Primary oculocerebral lymphoma	213812	Primitive neuroectodermal tumor of the cervix uteri	3235	Progressive deafness with stapes fixation
279897	Primary oculocerebral non-Hodgkin lymphoma	213630	Primitive neuroectodermal tumor of the corpus uteri	216812	Progressive deforming osteogenesis imperfecta
238606	Primary orthostatic tremor	854	Primitive portal vein thrombosis	217396	Progressive demyelinating neuropathy with bilateral striatal necrosis
439737	Primary PAN	3033	Primitive renal tubule syndrome		
99878	Primary parathyroid hyperplasia	2636	Primordial microcephalic dwarfism, Crachami type	1328	Progressive diaphyseal dysplasia
875	Primary pediatric cardiac tumor				
439737	Primary periarteritis nodosa	→2637	Primordial short stature-microdontia-opalescent and rootless teeth syndrome	495	Progressive diffuse palmoplantar keratoderma
168829	Primary peritoneal carcinoma				
168829	Primary peritoneal serous carcinoma	3042	Primrose syndrome	495	Progressive diffuse PPK
398980	Primary peritoneal serous/papillary carcinoma	397606	Prion protein systemic amyloidosis	438266	Progressive encephalomyelitis with rigidity and myoclonus
189439	Primary pigmented nodular adrenocortical disease	412066	PRKAR1B-related neurodegenerative dementia with intermediate filaments	2836	Progressive encephalopathy with edema, hypsarrhythmia and optic atrophy
100021	Primary plasmacytoma of the bone	2965	PRL-secreting pituitary adenoma	431361	Progressive encephalopathy with leukodystrophy due to DECR deficiency
439737	Primary polyarteritis nodosa	2965	PRLoma		
314566	Primary progressive apraxia of speech	326	Proaccelerin deficiency	99852	Progressive encephalopathy with severe infantile anorexia
75567	Primary progressive freezing gait	141099	Proboscis lateralis		
275766	Primary pulmonary arterial hypertension	740	Progeria	2836	Progressive encephalopathy-optic atrophy syndrome
2420	Primary pulmonary lymphoma	99706	Progeria-associated arthropathy		
358	Primary renal tubular hypokalemic hypomagnesemia with hypocalciuria	2959	Progeria-short stature-pigmented nevi syndrome	1947	Progressive epilepsy-intellectual disability syndrome, Finnish type
412206	Primary retention of teeth	300382	Progeroid and marfanoid aspect-lipodystrophy syndrome	457212	Progressive essential tremor-speech impairment-facial dysmorphism-intellectual disability-abnormal behavior syndrome
171	Primary sclerosing cholangitis	435953	Progeroid features-hepatocellular carcinoma predisposition syndrome		
99856	Primary syringomyelia	2962	Progeroid syndrome, De Barsy type	2744	Progressive external ophthalmoplegia and scoliosis
98841	Primary systemic ALCL	2963	Progeroid syndrome, Petty type		
314701	Primary systemic amyloidosis	79094	Progressive arterial occlusive disease-hypertension-heart defects-bone fragility-brachysyndactyly syndrome	352447	Progressive external ophthalmoplegia-myopathy-emaciation syndrome
268861	Primary tethered cord syndrome				
268861	Primary tethered spinal cord syndrome	448251	Progressive autosomal recessive ataxia-deafness syndrome	1214	Progressive facial hemiatrophy
99867	Primary thymic epithelial neoplasm				
263310	Primary thymic epithelial neoplasm type A	448251	Progressive autosomal recessive ataxia-sensorineural hearing loss syndrome	172	Progressive familial intrahepatic cholestasis
263324	Primary thymic epithelial neoplasm type AB				
263317	Primary thymic epithelial neoplasm type B	75373	Progressive bifocal chorioretinal atrophy	79306	Progressive familial intrahepatic cholestasis type 1
99867	Primary thymic epithelial tumor	→97229	Progressive bulbar palsy of childhood	79304	Progressive familial intrahepatic cholestasis type 2
263310	Primary thymic epithelial tumor type A	→97229	Progressive bulbar paralysis of childhood	79305	Progressive familial intrahepatic cholestasis type 3
263324	Primary thymic epithelial tumor type AB	139447	Progressive cavitating leukoencephalopathy	480483	Progressive familial intrahepatic cholestasis type 4
				480476	Progressive familial intrahepatic cholestasis type 5

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75327	Progressive foveal dystrophy	457265	Progressive myoclonus epilepsy type 9	742	Prolidase deficiency
1214	Progressive hemifacial atrophy			492	Proliferating trichilemmal cyst
199282	Progressive isolated segmental anhidrosis	352596	Progressive myoclonus epilepsy with dystonia	86872	Proliferation of large granular lymphocytes
73	Progressive massive osteolysis	726	Progressive neuronal degeneration of childhood with liver disease	221126	Proliferative vasculopathy and hydranencephaly/hydrocephaly
477814	Progressive microcephaly-seizures-cortical blindness-developmental delay syndrome			419	Proline oxidase deficiency
217260	Progressive multifocal leukoencephalitis	228012	Progressive neurosensory deafness-hypertrophic cardiomyopathy syndrome	75374	Prolonged electroretinal response suppression
217260	Progressive multifocal leukoencephalopathy			300878	Polymphocytic variant of hairy cell leukemia
454706	Progressive muscular atrophy	158022	Progressive nodular histiocytosis	300878	Polymphocytic variant of HCL
424027	Progressive myoclonic epilepsy due to CERS1 deficiency	100070	Progressive non-fluent aphasia	2083	Prominent glabella-microcephaly-hypogenitalism syndrome
263516	Progressive myoclonic epilepsy due to KCTD7 deficiency	2062	Progressive non-infectious anterior vertebral fusion	2966	Properdin deficiency
435438	Progressive myoclonic epilepsy due to KV3.1 deficiency	2762	Progressive osseous heteroplasia	35	Propionic acidemia
457265	Progressive myoclonic epilepsy due to LMNB2 deficiency			35	Propionic aciduria
308	Progressive myoclonic epilepsy type 1	3322	Progressive pancytopenia-immunodeficiency-cerebellar hypoplasia syndrome	35	Propionyl-CoA carboxylase deficiency
501	Progressive myoclonic epilepsy type 2	1159	Progressive pseudorheumatoid arthropathy of childhood	485358	Propylthiouracil embryofetopathy
263516	Progressive myoclonic epilepsy type 3	352718	Progressive retinal dystrophy due to retinol transport defect	485358	Propylthiouracil embryopathy
163696	Progressive myoclonic epilepsy type 4	447977	Progressive scapulohumeroperoneal distal myopathy	324977	Proteasome disability syndrome
402082	Progressive myoclonic epilepsy type 5	228012	Progressive sensorineural deafness-hypertrophic cardiomyopathy syndrome	324977	Proteasome-associated autoinflammatory syndrome
280620	Progressive myoclonic epilepsy type 6			213	Protein defect of cystin transport
435438	Progressive myoclonic epilepsy type 7	228012	Progressive sensorineural hearing loss-hypertrophic cardiomyopathy syndrome	26349	Protein S acquired deficiency
424027	Progressive myoclonic epilepsy type 8			744	Proteus syndrome
457265	Progressive myoclonic epilepsy type 9	457395	Progressive spondyloepimetaphyseal dysplasia-short stature-short fourth metatarsals-intellectual disability syndrome	2969	Proteus-like syndrome
352596	Progressive myoclonic epilepsy with dystonia			325	Prothrombin deficiency
308	Progressive myoclonus epilepsy type 1	683	Progressive supranuclear palsy	411696	Proton-pump inhibitor-responsive esophageal eosinophilia
501	Progressive myoclonus epilepsy type 2	240112	Progressive supranuclear palsy-apraxia of speech syndrome	251598	Protoplasmic astrocytoma
263516	Progressive myoclonus epilepsy type 3	240103	Progressive supranuclear palsy-corticobasal syndrome	79473	Protoporphyrinogen oxidase deficiency
402082	Progressive myoclonus epilepsy type 5	240085	Progressive supranuclear palsy-parkinsonism syndrome	2508	Proud syndrome
280620	Progressive myoclonus epilepsy type 6	240112	Progressive supranuclear palsy-progressive non-fluent aphasia syndrome	2508	Proud-Levine-Carpenter syndrome
435438	Progressive myoclonus epilepsy type 7			52022	Proximal 11p deletion syndrome
424027	Progressive myoclonus epilepsy type 8	240094	Progressive supranuclear palsy-pure akinesia with gait freezing syndrome	261197	Proximal 16p11.2 microdeletion syndrome
		316	Progressive symmetric erythrokeratoderma	370079	Proximal 16p11.2 microduplication syndrome
				261197	Proximal del(16)(p11.2)
		316	Progressive symmetric erythrokeratoderma, Gottron type	370079	Proximal dup(16)(p11.2)
				261197	Proximal monosomy 16p11.2
		2965	Prolactin-secreting pituitary adenoma	401768	Proximal myopathy with extrapyramidal signs
				606	Proximal myotonic dystrophy
		2965	Prolactinoma	606	Proximal myotonic myopathy
				47159	Proximal renal tubular acidosis
				93607	Proximal renal tubular acidosis with ocular abnormalities and intellectual disability
				70	Proximal spinal muscular atrophy
				83330	Proximal spinal muscular atrophy type 1

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
83418	Proximal spinal muscular atrophy type 2	88939	Pseudohypoadosteronism type 2B	240112	PSP-PNFA
		88940	Pseudohypoadosteronism type 2C	240094	PSP-pure akinesia with gait freezing
83419	Proximal spinal muscular atrophy type 3	300525	Pseudohypoadosteronism type 2D	263548	PSS type A
		300530	Pseudohypoadosteronism type 2E	263553	PSS type B
83420	Proximal spinal muscular atrophy type 4	79443	Pseudohypoparathyroidism type 1A	99928	PSST
		94089	Pseudohypoparathyroidism type 1B	71519	Psychogenic dystonia
3250	Proximal symphalangism	79444	Pseudohypoparathyroidism type 1C	71519	Psychogenic movement disorders
370079	Proximal trisomy 16p11.2	94090	Pseudohypoparathyroidism type 2	324636	Psychogenic purpura
3390	Proximal tubulopathy-diabetes mellitus-cerebellar ataxia syndrome	2976	Pseudoprechaunism syndrome, Patterson type	88618	Psychomotor retardation due to S-adenosylhomocysteine hydrolase deficiency
397606	PrP systemic amyloidosis	26790	Pseudomyxoma peritonei		
3222	PRPP synthetase superactivity			52530	PT-VWD
3222	PRPS1 superactivity	251962	Pseudopapillary ganglioglioneurocytoma	329	PTA deficiency
47159	pRTA			247698	PTC syndrome
2970	Prune belly syndrome	251962	Pseudopapillary neurocytoma with glial differentiation	97290	PTC-RCC
89843	Pruriginous dystrophic epidermolysis bullosa			269229	PTCD
64745	Pruritic urticarial papules and plaques of pregnancy	2980	Pseudopapilledema-blepharophimosis-hand anomalies syndrome	2988	Pterygium colli-intellectual disability-digital anomalies syndrome
284417	PSAT deficiency	129	Pseudopelade of Brocq		
171	PSC	2985	Pseudoprogeria syndrome	86789	PTLAH
228402	Pseudo-Angelman syndrome	79445	Pseudopseudohypoparathyroidism	70568	PTLD
99000	Pseudo-Best disease	477742	Pseudosarcomatous fasciitis		
314459	Pseudo-Demons-Meigs syndrome	477742	Pseudosarcomatous fibromatosis	2999	Ptosis-strabismus-ectopic pupils syndrome
577	Pseudo-Hurler polydystrophy	3103	Pseudothalidomide syndrome	→293843	Ptosis-strabismus-rectus abdominis diastasis syndrome
314459	Pseudo-Meigs syndrome	2518	Pseudotoxoplasmosis syndrome		
439881	Pseudo-membranous bronchitis	238624	Pseudotumor cerebri	238766	Ptosis-syndactyly-learning difficulties syndrome
263482	Pseudo-Morquio syndrome type 2	83316	Pseudotymphus of California		
2971	Pseudo-NALD	180079	Pseudounicornuate uterus		
2971	Pseudo-neonatal adrenoleukodystrophy	753	Pseudovaginal perineoscrotal hypospadias	228396	Ptosis-upper ocular movement limitation-absence of lacrimal punctum syndrome
1229	Pseudo-TORCH syndrome	289157	Pseudovitamin D-deficient rickets	2997	Ptosis-vocal cord paralysis syndrome
2166	Pseudo-trisomy 13 syndrome	758	Pseudoxanthoma elasticum	251915	PTPR
99000	Pseudo-vitelliform macular dystrophy	228293	Pseudoxanthoma elasticum-like papillary dermal elastolysis	485358	PTU embryofetopathy
52530	Pseudo-von Willebrand disease			485358	PTU embryopathy
52530	Pseudo-von Willebrand disease type 2B	436274	Pseudoxanthoma elasticum-like skin manifestations with retinitis pigmentosa	231580	PUAH
→300	Pseudo-Zellweger syndrome			60039	Pudendal algia
750	Pseudoachondroplasia	91135	Pseudoxanthoma elasticum-like syndrome	60039	Pudendal nerve entrapment syndrome
750	Pseudoachondroplastic dysplasia	228227	Pseudoxanthoma-like late-onset focal dermal elastosis	60039	Pudendal neuralgia
750	Pseudoachondroplastic spondyloepiphyseal dysplasia	280794	Pseudoxanthomatous DCM	60039	Pudendal neuralgia by pudendal nerve entrapment
2971	Pseudoachondroplastic spondyloepiphyseal dysplasia	280794	Pseudoxanthomatous diffuse cutaneous mastocytosis	60039	Pudendal algia
526	Pseudoaldosteronism	95496	PSIS	443173	Puerperal psychosis
221120	Pseudoaminopterin syndrome	85436	Psoriasis-related JIA	984	Pulmonary agenesis
85174	Pseudodiastrophic dysplasia	85436	Psoriasis-related juvenile idiopathic arthritis	60025	Pulmonary alveolar microlithiasis
2983	Pseudohermaphroditism-intellectual disability syndrome	683	PSP syndrome	440427	Pulmonary alveolar proteinosis, Reunion island type
526	Pseudohyperaldosteronism type 1	240112	PSP-AOS		
88660	Pseudohyperaldosteronism type 2	240103	PSP-CBS	→331176	Pulmonary arterial hypertension-leukopenia-atrial septal defect syndrome
756	Pseudohypoadosteronism type 1	240103	PSP-corticobasal syndrome		
757	Pseudohypoadosteronism type 2	240085	PSP-p	2038	Pulmonary arteriovenous malformation
88938	Pseudohypoadosteronism type 2A	240094	PSP-PAGF		
		240085	PSP-parkinsonism		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
99049	Pulmonary artery coming from patent ductus arteriosus	308013	Punctate palmoplantar keratoderma type 3 without elastoidosis	3005	Pyle disease
99050	Pulmonary artery coming from the aorta			48104	Pyoderma gangrenosum
99083	Pulmonary artery hypoplasia	438213	PURA-related severe neonatal hypotonia-seizures-encephalopathy syndrome	289478	Pyoderma gangrenosum-acne-suppurative hidradenitis syndrome
1207	Pulmonary atresia with ventricular septal defect			69126	Pyogenic arthritis-pyoderma gangrenosum-acne syndrome
1208	Pulmonary atresia-intact ventricular septum syndrome	438216	PURA-related severe neonatal hypotonia-seizures-encephalopathy syndrome due to a point mutation	183713	Pyogenic bacterial infections due to MyD88 deficiency
64741	Pulmonary blastoma			764	Pyomyositis
99084	Pulmonary branch stenosis	231625	Pure aldosterone-producing adrenocortical carcinoma	2561	Pyramidal molar-glaucoma-upper abnormal lip syndrome
199241	Pulmonary capillary hemangiomas			63440	Pyrgocephaly
210136	Pulmonary fibrosis-hepatic hyperplasia-bone marrow hypoplasia syndrome	231625	Pure APAC	79096	Pyridoxal phosphate-dependent seizures
		441	Pure autonomic failure	79096	Pyridoxal phosphate-responsive seizures
		441	Pure dysautonomia		
217080	Pulmonary fungal infections in patients deemed at risk	319465	Pure familial acute myeloid leukemia	79096	Pyridoxamine 5'-oxidase deficiency
99874	Pulmonary histiocytosis X	319465	Pure familial AML	79096	Pyridoxamine 5'-phosphate oxidase deficiency
991	Pulmonary hypoplasia-agonadism-dextrocardia-diaphragmatic hernia syndrome	69084	Pure hair and nail ectodermal dysplasia	3006	Pyridoxine-dependent epilepsy
		441	Pure idiopathic dysautonomia	32	Pyroglutamic aciduria
217557	Pulmonary interstitial glycogenosis	475	Pure Joubert syndrome	293633	Pyroline-5-carboxylate reductase 1 deficiency
2414	Pulmonary lymphangiomatosis	254854	Pure mitochondrial myopathy	3008	Pyruvate carboxylase deficiency
60026	Pulmonary nodular lymphoid hyperplasia	2028	Puretic syndrome	353308	Pyruvate carboxylase deficiency type A
411703	Pulmonary non-tuberculous mycobacterial infection	760	Purine nucleoside phosphorylase deficiency		
		761	Purpura rheumatica	353314	Pyruvate carboxylase deficiency type B
60026	Pulmonary pseudolymphoma	2442	Purtilo syndrome	353320	Pyruvate carboxylase deficiency type C
101206	Pulmonary valve agenesis-tetralogy of Fallot-absence of ductus arteriosus syndrome	293173	Pustular drug eruption	353320	Pyruvate carboxylase deficiency, benign type
		163927	Pustulosis palmaris et plantaris		
		48377	Pustulosis subcornealis		
99048	Pulmonary valve agenesis-ventricular septal defect-persistent ductus arteriosus syndrome	93110	PUV	353308	Pyruvate carboxylase deficiency, infantile form
31837	Pulmonary venoocclusive disease	729	PV	353314	Pyruvate carboxylase deficiency, severe neonatal type
		101206	PVA/ADA, Fallot type		
→636	Pulmonic stenosis with 'café-au-lait' spots	99048	PVA/PDA, non-Fallot type	79243	Pyruvate decarboxylase deficiency
85202	Pulmonic stenosis-brachytelephalangism-calcification of cartilages syndrome	398069	PWS due to point mutation	79244	Pyruvate dehydrogenase complex component E2 deficiency
		398073	PWS-like		
		398079	PWS-like due to point mutation	255182	Pyruvate dehydrogenase complex component E3 deficiency
98984	Pulverulent cataract	251607	PXA	765	Pyruvate dehydrogenase complex deficiency
97353	Punch-drunk syndrome	758	PXE		
79502	Punctate palmoplantar hyperkeratosis type 2	228227	PXE-like late-onset focal dermal elastosis	79243	Pyruvate dehydrogenase complex E1 component subunit alpha deficiency
		228293	PXE-like papillary dermal elastolysis		
38	Punctate palmoplantar hyperkeratosis type 3	91135	PXE-like syndrome	255138	Pyruvate dehydrogenase complex E1 component subunit beta deficiency
		436274	PXE-like syndrome with retinitis pigmentosa		
308013	Punctate palmoplantar hyperkeratosis type 3 without elastoidosis	763	Pycnodysostosis	765	Pyruvate dehydrogenase deficiency
		293633	PYCR1 deficiency		
79501	Punctate palmoplantar keratoderma type 1	293633	PYCR1-related De Barsy syndrome	79243	Pyruvate dehydrogenase E1-alpha deficiency
79502	Punctate palmoplantar keratoderma type 2	481152	PYCR2-related microcephaly-progressive leukoencephalopathy		
		3003	Pyknoachondrogenesis	255138	Pyruvate dehydrogenase E1-beta deficiency
38	Punctate palmoplantar keratoderma type 3	763	Pyknodysostosis	79244	Pyruvate dehydrogenase E2 deficiency
		64280	Pyknolepsy		

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2394	Pyruvate dehydrogenase E3 deficiency	295219	Radio-ulnar fusion, bilateral	→1071	Rapp-Hodgkin syndrome
		295217	Radio-ulnar fusion, unilateral	213528	Rare adenocarcinoma of the breast
255182	Pyruvate dehydrogenase E3-binding protein deficiency	295219	Radio-ulnar synostosis, bilateral	137820	Rare endometriosis
		295217	Radio-ulnar synostosis, unilateral	98345	Rare idiopathic male infertility
79246	Pyruvate dehydrogenase phosphatase deficiency	71289	Radio-ulnar synostosis-amegakaryocytic thrombocytopenia syndrome	98619	Rare isolated myopia
255182	Pyruvate dehydrogenase protein X component deficiency			101685	Rare non-syndromic intellectual disability
766	Pyruvate kinase deficiency of erythrocytes	→193	Radio-ulnar synostosis-retinal pigment abnormalities syndrome	101685	Rare NSID
781	Q fever	294979	Radio-ulnar terminal transverse meromelia	213574	Rare variants of adenocarcinoma of the corpus uteri
3010	Qazi-Markouizos syndrome			75564	RARS
602	Quadriceps-sparing myopathy	295095	Radio-ulnar terminal transverse meromelia, bilateral	438114	RARS-related autosomal recessive hypomyelinating leukodystrophy
781	Quadrilateral fever	295093	Radio-ulnar terminal transverse meromelia, unilateral	268114	RAS-associated autoimmune leukoproliferative disease
9	Quadruple X				
84142	Quantal squander syndrome	420741	Radiosensitivity-immunodeficiency-dysmorphic features-learning difficulties syndrome	1929	Rasmussen subacute encephalitis
869	Quaternary A syndrome			1929	Rasmussen syndrome
220436	Quebec platelet disorder	3269	Radioulnar fusion	3023	Rasmussen-Johnsen-Thomsen syndrome
781	Query fever				
137888	Question mark ear syndrome	3270	Radioulnar synostosis-developmental delay-hypotonia syndrome	31205	Rat-bite fever
346	Quinquaud's folliculitis decalvans			436	Rathburn disease
261529	r(Y)			99852	Ravine syndrome
100057	RAAS-blocker-induced angioedema	3268	Radioulnar synostosis-microcephaly-scoliosis syndrome	2840	Ray-Peterson-Scott syndrome
100057	RAAS-blocker-induced angioneurotic edema	100057	RAE	79127	RB-ILD
770	Rabies	100019	RAEB-1	98961	RBCD
769	Rabson-Mendenhall syndrome	100020	RAEB-2	93111	RCAD syndrome
240760	RAD50 deficiency	168960	RAEB-t	177	RCDP
93321	Radial clubhand	1832	Raine syndrome	284388	RCVS
		50811	Rajab-Spranger syndrome	79408	RDEB generalisata gravis
1121	Radial deficiency-tibial hypoplasia syndrome	268114	RALD	89842	RDEB generalisata mitis
93321	Radial hemimelia	99843	Rambam-Hasharon syndrome	89841	RDEB, centripetalis
295071	Radial hemimelia, bilateral	3018	Rambaud-Gallian syndrome	79408	RDEB, Hallopeau-Siemens type
295069	Radial hemimelia, unilateral	3018	Rambaud-Gallian-Touchard syndrome	89842	RDEB, non-Hallopeau-Siemens type
		3019	Ramon syndrome	89841	RDEB-Ce
2252	Radial hypoplasia-triphalangeal thumbs-hypospadias-maxillary diastema syndrome	1051	Ramos-Arroyo syndrome	89842	RDEB-generalized other
		3020	Ramsay Hunt syndrome	79409	RDEB-I
93321	Radial longitudinal meromelia	86861	Randall disease	89842	RDEB-O
295071	Radial longitudinal meromelia, bilateral	3021	RAPADILINO syndrome	79408	RDEB-sev gen
		293987	Rapid-onset childhood obesity-hypothalamic dysfunction-hypoventilation-autonomic dysregulation syndrome	85445	Reactive amyloidosis
295069	Radial longitudinal meromelia, unilateral			29207	Reactive arthritis
93321	Radial ray agenesis			314962	Reactive hypereosinophilic syndrome
		293987	Rapid-onset childhood obesity-hypothalamic dysfunction-hypoventilation-autonomic dysregulation-neural tumors syndrome	166433	Reading seizures
2307	Radial ray defects, hearing impairment, external ophthalmoplegia, and thrombocytopenia			857	REAR syndrome
				1188	Reardon-Baraitser syndrome
3026	Radial ray hypoplasia-choanal atresia syndrome			2631	Reardon-Hall-Slaney syndrome
90021	Radiation myelitis	71517	Rapid-onset dystonia-parkinsonism	96167	Rec8 syndrome
70475	Radiation proctitis	141184	Rapidly involuting congenital hemangioma	96167	Rec(8) syndrome
99789	Radicular dentin dysplasia	280569	Rapidly progressive glomerulonephritis	1115	Recessive aplasia cutis congenita of limbs
→2712	Radiculomegaly of canine teeth-congenital cataract			79409	Recessive dystrophic epidermolysis bullosa inversa
3015	Radio-renal syndrome	178307	RAPK		

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89842	Recessive dystrophic epidermolysis bullosa, non-Hallopeau-Siemens type	98826	Refractory anemia	93111	Renal dysfunction-early-onset diabetes syndrome
		86839	Refractory anemia with excess blasts	93108	Renal dysplasia
89842	Recessive dystrophic epidermolysis bullosa-generalized other	168960	Refractory anemia with excess blasts in transformation	93173	Renal dysplasia, bilateral
				93172	Renal dysplasia, unilateral
280384	Recessive intellectual disability-motor dysfunction-multiple joint contractures syndrome	100019	Refractory anemia with excess blasts type 1	3404	Renal dysplasia-limb defects syndrome
94125	Recessive mitochondrial ataxia syndrome	100020	Refractory anemia with excess blasts type 2	→1768	Renal dysplasia-megalocystis-sirenomelia syndrome
461	Recessive X-linked ichthyosis	75564	Refractory anemia with ringed sideroblasts	3404	Renal dysplasia-mesomelia-radiohumeral fusion syndrome
96167	Recombinant 8 syndrome	398063	Refractory CD	3156	Renal dysplasia-retinal aplasia syndrome
96167	Recombinant chromosome 8 syndrome	398063	Refractory celiac disease		
99990	Recrudescence typhus	398063	Refractory sprue	140969	Renal dysplasia-retinal pigmentary dystrophy-cerebellar ataxia-skeletal dysplasia syndrome
171220	Rectal duplication	773	Refsum disease		
100081	Rectal neuroendocrine tumor	1525	Reginato-Schiapachasse syndrome	654	Renal embryonic tumor
424002	Rectal squamous cell carcinoma	1433	Regional choroidal atrophy and alopecia	1652	Renal Fanconi syndrome with nephrocalcinosis and renal stones
51890	Rectus abdominis syndrome	83450	Regional odontodysplasia	34528	Renal hypomagnesemia type 2
88619	Recurrent acute necrotizing encephalopathy	300865	Regressive atypical histiocytosis	31043	Renal hypomagnesemia type 3
64740	Recurrent acute pancreatitis	1040	Regressive metaphyseal dysplasia	93101	Renal hypoplasia
		448267	Regressive spondylometaphyseal dysplasia	97362	Renal hypoplasia, bilateral
2672	Recurrent encephalopathy of childhood	2634	Reinhardt-Pfeiffer mesomelic dysplasia	97361	Renal hypoplasia, unilateral
90052	Recurrent hepatitis C virus induced liver disease in liver transplant recipients	2634	Reinhardt-Pfeiffer syndrome	319319	Renal medullary carcinoma
		98961	Reis-Bücklers corneal dystrophy	71273	Renal nutcracker syndrome
293381	Recurrent hereditary corneal erosions	29207	Reiter disease	171871	Renal pseudohypoadosteronism type 1
		29207	Reiter syndrome	18	Renal tubular acidosis type 1
169142	Recurrent infection due to specific granule deficiency	99991	Relapsing epidemic typhus	47159	Renal tubular acidosis type 2
183675	Recurrent infections associated with rare immunoglobulin isotypes deficiency	33577	Relapsing febrile nodular non-suppurative panniculitis	2785	Renal tubular acidosis type 3
		33577	Relapsing febrile nodular panniculitis	3033	Renal tubular dysgenesis
251523	Recurrent infections-inflammatory syndrome due to zinc metabolism disorder syndrome	91547	Relapsing fever	97369	Renal tubular dysgenesis due to twin-twin transfusion
		728	Relapsing polychondritis	97369	Renal tubular dysgenesis of genetic origin
69665	Recurrent intrahepatic cholestasis of pregnancy	412	Remnant disease	112	Renal tubular normotensive hypokalemic alkalosis with hypercalciuria
		217330	REN-associated familial juvenile hyperuricemic nephropathy		
480864	Recurrent metabolic encephalomyopathic crises-rhabdomyolysis-cardiac arrhythmia-intellectual disability syndrome	217330	REN-associated FJHN	254902	Renal tubulopathy-encephalopathy-liver failure syndrome
		217330	REN-associated kidney disease	857	Renal-ear-anal-radial syndrome
		411709	Renal agenesis	1092	Renal-genital-middle ear anomalies
169467	Recurrent Neisseria infections due to factor D deficiency	1848	Renal agenesis, bilateral	294415	Renal-hepatic-pancreatic dysplasia
		93100	Renal agenesis, unilateral	3032	Renal-hepatic-pancreatic dysplasia-Dandy-Walker cysts syndrome
60032	Recurrent respiratory papillomatosis	2838	Renal caliceal diverticuli-deafness syndrome	774	Rendu-Osler disease
199267	Recurring digital fibrous tumor of childhood	319314	Renal cell carcinoma after neuroblastoma	774	Rendu-Osler-Weber disease
79433	Red oculocutaneous albinism	319314	Renal cell carcinoma associated with neuroblastoma	93975	Renier-Gabreels-Jasper syndrome
231031	Red palms disease			100057	Renin-angiotensin-aldosterone system-blocker-induced angioedema
838	RED-M	1475	Renal coloboma syndrome	100057	Renin-angiotensin-aldosterone system-blocker-induced angioneurotic edema
97239	Reducing body myopathy	93111	Renal cysts and diabetes syndrome		
523	Reed syndrome	93111	Renal cysts-maturity-onset diabetes of the young syndrome	294415	Renohepaticpancreatic dysplasia
3221	Refetoff syndrome				
99995	Reflex sympathetic dystrophy				

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3033	Renotubular dysgenesis	85332	Retinitis pigmentosa and intellectual disability due to Xp11.3 microdeletion	69077	Rhabdoid tumor
3242	Renpenning syndrome			231108	Rhabdoid tumor predisposition syndrome
73273	Resistance to IGF-1			3097	Rhabdomyomatous dysplasia-cardiopathy-genital anomalies syndrome
424	Resistance to thyroid stimulating hormone				
99832	Resistance to thyrotropin-releasing hormone syndrome	140976	Retinitis pigmentosa-hypopituitarism-nephronophthisis-skeletal dysplasia syndrome	780	Rhabdomyosarcoma
247257	Respiratory anthrax	3085	Retinitis pigmentosa-intellectual disability-deafness-hypogenitalism syndrome	213802	Rhabdomyosarcoma of the cervix uteri
247257	Respiratory anthrax disease			213615	Rhabdomyosarcoma of the corpus uteri
79127	Respiratory bronchiolitis-interstitial lung disease syndrome			3099	Rheumatic fever
1662	Restrictive dermopathy	436245	Retinitis pigmentosa-juvenile cataract-short stature-intellectual disability syndrome	85408	Rheumatoid factor-negative JIA
33355	Reticular dysgenesis			247854	Rheumatoid factor-negative JIA with anti-nuclear antibodies
99002	Reticular dystrophy of the retinal pigment epithelium	52427	Retinitis punctata albescens	247861	Rheumatoid factor-negative JIA without anti-nuclear antibodies
100000	Reticular perineurioma	790	Retinoblastoma		
79145	Reticular pigment anomaly of flexures	838	Retinocochleocerebral vasculopathy	85408	Rheumatoid factor-negative juvenile idiopathic arthritis
178307	Reticulate acropigmentation of Kitamura	3087	Retinohepatoendocrinologic syndrome		
86900	Reticulum cell sarcoma	2305	Retinoic acid embryopathy	247854	Rheumatoid factor-negative juvenile idiopathic arthritis with anti-nuclear antibodies
458763	Retiform hemangioendothelioma	40366	Retinoid embryopathy		
284247	Retinal arterial macroaneurysm and supraulvalvar pulmonic stenosis	2305	Retinoids embryopathy		
75326	Retinal arterial tortuosity	352718	Retinol dystrophy-iris coloboma-comedogenic acne syndrome	247861	Rheumatoid factor-negative juvenile idiopathic arthritis without anti-nuclear antibodies
75326	Retinal arteriolar tortuosity	90050	Retinopathy of prematurity		
71213	Retinal capillary malformation	139455	Retinopathy, Burgess-Black type	85435	Rheumatoid factor-positive polyarticular JIA
1574	Retinal degeneration-nanophthalmos-glaucoma syndrome	3088	Retinopathy-anemia-central nervous system anomalies syndrome		
1571	Retinal detachment-occipital encephalocele syndrome	838	Retinopathy-encephalopathy-deafness associated with microangiopathy	85435	Rheumatoid factor-positive polyarticular juvenile idiopathic arthritis
397758	Retinal dystrophy with inner nuclear layer and ganglion cell anomalies				
397758	Retinal dystrophy with inner retinal dysfunction and ganglion cell anomalies	53540	Retinoschisis with early nyctalopia	177	Rhizomelic chondrodysplasia punctata
436245	Retinal dystrophy-juvenile cataract-short stature syndrome	269200	Retrocerebellar cyst		
75326	Retinal hemorrhage with vascular tortuosity	90050	Retrolental fibroplasia	309789	Rhizomelic chondrodysplasia punctata type 1
3018	Retinal ischemic syndrome-digestive tract small vessel hyalinosis-diffuse cerebral calcifications syndrome	778	Rett syndrome		
319640	Retinal macular dystrophy type 2	3095	Rett syndrome variant	309796	Rhizomelic chondrodysplasia punctata type 2
247691	Retinal vasculopathy and cerebral leukoencephalopathy	294049	Reunion Island Larsen syndrome		
353356	Retinal vasoproliferative tumor	99852	Reunion island-anorexia-vomiting which is irrepressible-neurological signs syndrome	309803	Rhizomelic chondrodysplasia punctata type 3
791	Retinitis pigmentosa			468717	Rhizomelic chondrodysplasia punctata type 5
85332	Retinitis pigmentosa and intellectual disability due to del(X)(p11.3)	284388	Reversible cerebral vasoconstriction syndrome	2831	Rhizomelic dysplasia, Patterson-Lowry type
85332	Retinitis pigmentosa and intellectual disability due to monosomy Xp11.3	254864	Reversible infantile cytochrome C oxidase deficiency		
		254864	Reversible infantile respiratory chain deficiency	93569	Rhizomelic pseudopolyarthritis
		3088	Revesz syndrome	1453	Rhizomelic shortness with clavicular defect
		3088	Revesz-DeBuse syndrome		
		3096	Reye syndrome	3098	Rhizomelic syndrome, Urbach type
		199267	Reye tumor	59315	Rhombencephalosynapsis
		779	Reynolds syndrome	→1071	RHS
		244310	RFT1-CDG	140976	RHYNS syndrome
		251975	RGNT	217055	RI-CMT type A
		71275	Rh deficiency syndrome	254334	RI-CMT type B
		71275	Rh-null syndrome	369867	RI-CMT type C
				435998	RI-CMT type D
				97229	Riboflavin transporter deficiency

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
440706	Ribose-5-P isomerase deficiency	96173	Ring chromosome 9	1945	Rolandic epilepsy
141184	RICH	1438	Ring chromosome 10	163727	Rolandic epilepsy-paroxysmal exercise-induced dystonia-writer's cramp syndrome
1399	Richards-Rundle syndrome	96175	Ring chromosome 11	163721	Rolandic epilepsy-speech dyspraxia syndrome
240071	Richardson syndrome	1439	Ring chromosome 12	101016	Romano-Ward long QT syndrome
2323	Richardson-Kirk syndrome	96176	Ring chromosome 13	101016	Romano-Ward syndrome
3101	Richieri Costa-da Silva syndrome	1440	Ring chromosome 14	1214	Romberg syndrome
→2995	Richieri Costa-Guion Almeida syndrome	96177	Ring chromosome 15	3110	Rombo syndrome
		96178	Ring chromosome 16	1088	Rommen-Mueller-Sybert syndrome
2511	Richieri Costa-Guion Almeida-Ramos syndrome	1441	Ring chromosome 17	90050	ROP
		1442	Ring chromosome 18	158014	Rosaï-Dorfman disease
→2353	Richieri Costa-Guion Almeida-Rodini syndrome	1443	Ring chromosome 19	158014	Rosaï-Dorfman-Destombes disease
		1444	Ring chromosome 20	1837	Rosenberg-Lohr syndrome
3102	Richieri Costa-Pereira syndrome	1445	Ring chromosome 21	329	Rosenthal factor deficiency
1784	Richieri-Costa-Colletto syndrome	1446	Ring chromosome 22	329	Rosenthal syndrome
1794	Richieri-Costa-Gorlin syndrome	261529	Ring chromosome Y	251975	Rosette-forming glioneuronal tumor of fourth ventricle
28378	Richner-Hanhart syndrome	91481	Ring dermoid of cornea	2909	Rothmund-Thomson syndrome
606	Ricker disease	91481	Ring dermoid syndrome	221008	Rothmund-Thomson syndrome type 1
606	Ricker syndrome	169	Ringed hair disease	221016	Rothmund-Thomson syndrome type 2
83312	Rickettsialpox	97238	Rippling muscle disease	3111	Rotor syndrome
420741	RIDDLE syndrome	206575	Rippling muscle disease with myasthenia gravis	3115	Roussy-Lévy syndrome
64744	Riedel disease	7	Ritscher-Schinzel syndrome	1323	Rozin-camptodactyly syndrome
64744	Riedel thyroiditis	1803	Rivera-Perez-Salas syndrome	1323	Rozin-Hertz-Goodman syndrome
91483	Rieger anomaly	294049	RLS	280569	RPGN
3163	Rieger anomaly-partial lipodystrophy syndrome	93307	rMED	1507	RRS
782	Rieger syndrome	137634	RNF135-related overgrowth syndrome	818	RSH syndrome
319251	Rift valley fever	420741	RNF168 deficiency	293848	RTLA
99081	Right aortic arch	71273	RNS	231108	RTPS
99119	Right inferior caval vein connecting to left-sided atrium	3103	Roberts syndrome	2909	RTS
99119	Right inferior vena cava connecting to left-sided atrium	3103	Roberts-SC phocomelia syndrome	221008	RTS1
99119	Right IVC connecting to left-sided atrium	3104	Robin sequence-oligodactyly syndrome	221016	RTS2
99110	Right superior caval vein connecting to left-sided atrium	97360	Robinow dwarfism	83616	Rubella panencephalitis
99110	Right superior vena cava connecting to left-sided atrium	97360	Robinow syndrome	783	Rubinstein-Taybi syndrome
99110	Right superior vena cava connecting to left-sided atrium	3105	Robinow-like syndrome	353281	Rubinstein-Taybi syndrome due to 16p13.3 microdeletion
99110	Right SVC connecting to left-sided atrium	97360	Robinow-Silverman-Smith syndrome	353277	Rubinstein-Taybi syndrome due to CREBBP mutations
97244	Rigid spine congenital muscular dystrophy	→794	Robinow-Sorauf syndrome	353284	Rubinstein-Taybi syndrome due to EP300 haploinsufficiency
97244	Rigid spine syndrome	2780	Robinow-Unger syndrome	1768	Rudd-Klimek syndrome
1764	Riley-Day syndrome	529	Roch-Leri mesosomatous lipomatosis	→798	Rudiger syndrome
217335	RIN2 deficiency	83311	Rocky Mountain spotted fever	79433	Rufous oculocutaneous albinism
217335	RIN2 syndrome	49382	Rod monochromacy	435953	Ruijs-Aalfs syndrome
1437	Ring chromosome 1	49382	Rod monochromatism	1672	Russell diencephalic cachexia
96171	Ring chromosome 2	1258	Rodini-Richieri Costa syndrome	1672	Russell syndrome
96172	Ring chromosome 3	49827	Rogers syndrome	1834	Russell-Weaver-Bull syndrome
1447	Ring chromosome 4	293987	ROHHAD	2709	Rutherford syndrome
251043	Ring chromosome 5	293987	ROHHADNET	3121	Ruvalcaba syndrome
1448	Ring chromosome 6	353298	Roifman syndrome	247691	RVCL
1449	Ring chromosome 7	221139	Roifman-Chitayat syndrome		
1450	Ring chromosome 8	→1855	Roifman-Melamed syndrome		
		247775	Rokitansky sequence		
		3109	Rokitansky syndrome		

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293848	rvFTD	70595	SANDO	→98769	SCA16
461	RXLI	2378	Sandrow syndrome	98759	SCA17
16	S cone monochromacy	581	Sanfilippo disease	98771	SCA18
16	S cone monochromatism	79269	Sanfilippo syndrome type A	98772	SCA19/22
3105	Saal-Greenstein syndrome	79270	Sanfilippo syndrome type B	101110	SCA20
319239	Sabia hemorrhagic fever	79271	Sanfilippo syndrome type C	98773	SCA21
3124	Saccharopine dehydrogenase deficiency	79272	Sanfilippo syndrome type D	→98772	SCA22
		2323	Sanjad-Sakati syndrome	101108	SCA23
3124	Saccharopinuria	588	Santavuori congenital muscular dystrophy	101111	SCA25
286	Sack-Barabas syndrome			101112	SCA26
98841	sACL	79263	Santavuori disease	98764	SCA27
3027	Sacral agenesis syndrome	79263	Santavuori-Haltia disease	101109	SCA28
397927	Sacral agenesis-abnormal ossification of the vertebral bodies-persistent notochordal canal syndrome	2155	Santos-Mateus-Leal syndrome	208513	SCA29
		98868	SAO	211017	SCA30
		247234	SAOA	217012	SCA31
		793	SAPHO syndrome	276183	SCA32
→83628	Sacral hemangiomas-multiple congenital abnormalities syndrome	54368	Sarcocystosis	1955	SCA34
		797	Sarcoidosis	276193	SCA35
2351	Sacral meningocele-conotruncal heart defects syndrome	3129	Sarcosine dehydrogenase complex deficiency	276198	SCA36
3027	Sacral regression syndrome	3129	Sarcosinemia	363710	SCA37
1773	Sacrococcygeal dysgenesis association	54368	Sarcosporidiosis	423296	SCA38
85165	SADDAN	1878	Sarcotubular myopathy	423275	SCA40
794	Saethre-Chotzen syndrome	3130	Satoyoshi syndrome	458798	SCA41
2872	Sagittal craniostenosis with congenital heart disease, mental deficiency and mandibular ankylosis	330015	Saturnism	458803	SCA42
		425120	SAVI	26792	SCAD deficiency
		3047	Say-Barber-Biesecker-Young-Simpson syndrome	26792	SCADD
300493	Saglikler syndrome			254881	SCAE
83484	Saint Louis encephalitis	2013	Say-Barber-Hobbs syndrome	1003	Scalp defects-postaxial polydactyly syndrome
2256	Saito-Kuba-Tsuruta syndrome	3132	Say-Barber-Miller syndrome	370052	SCALP syndrome
1409	Salamon syndrome	3133	Say-Field-Coldwell syndrome	2036	Scalp-ear-nipple syndrome
2613	Salcedo syndrome	3369	Say-Meyer syndrome	64753	SCAN 2
140969	Saldino-Mainzer syndrome	3047	SBBYSS	94124	SCAN1
213557	Salivary gland type cancer of the breast	79157	SBCAD deficiency	840	SCAP
213557	Salivary gland type carcinoma of the breast	481	SBMA	168624	Scaphocephaly-macrocephaly-maxillary retrusion-intellectual disability syndrome
		3103	SC phocomelia		
		3103	SC pseudothalidomide syndrome		
309334	Salla disease	98755	SCA1	2839	Scapuloiliac dysostosis
370938	Salt-and-pepper syndrome	98756	SCA2	431255	Scapulooperoneal neuronopathy
112	Salt-losing tubular disorder, Henle's loop type	98757	SCA3	431255	Scapulooperoneal spinal muscular atrophy
		276238	SCA3, Joseph type		
112	Salt-wasting tubulopathy, Henle's loop type	276244	SCA3, Machado type	64753	SCAR1
		276241	SCA3, Thomas type	1170	SCAR2
2230	Salti-Salem syndrome	98765	SCA4	95433	SCAR3
369992	SAM syndrome	98766	SCA5	95434	SCAR4
53721	SAMS 1-31	98758	SCA6	83472	SCAR5
397623	SAMS syndrome	94147	SCA7	284332	SCAR6
228123	San Joaquin valley fever	98760	SCA8	284324	SCAR7
96167	San Luis Valley syndrome	98761	SCA10	88644	SCAR8
796	Sandhoff disease	98767	SCA11	139485	SCAR9
309169	Sandhoff disease, adult form	98762	SCA12	284289	SCAR10
309155	Sandhoff disease, infantile form	98768	SCA13	284271	SCAR11
309162	Sandhoff disease, juvenile form	98763	SCA14	284282	SCAR12
71272	Sandifer syndrome	98769	SCA15/16	324262	SCAR13

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352403	SCAR14	50944	Schöpf-Schulz-Passarge syndrome	295193	SD1, Castilla type
404499	SCAR15	93921	Schwannomatosis	295189	SD1, Lueken type
412057	SCAR16	800	Schwartz-Jampel syndrome	295191	SD1, Montagu type
453521	SCAR17	800	Schwartz-Jampel syndrome type 1	295187	SD1, Weidenreich type
363432	SCAR18	3206	Schwartz-Jampel syndrome type 2	295187	SD1a
448251	SCAR19	800	Schwartz-Jampel-Aberfeld syndrome	295189	SD1b
397709	SCAR20			295191	SD1c
466794	SCAR21	277	SCID due to adenosine deaminase deficiency	295193	SD1d
404493	SCAR23			295197	SD2, Debeer type
3134	SCARF syndrome	275	SCID due to ARTEMIS deficiency	295199	SD2, Malik type
90080	Scarring in glaucoma filtration surgical procedures	357237	SCID due to CARD11 deficiency	295195	SD2, Vordingborg type
		331206	SCID due to complete RAG1/2 deficiency	295195	SD2a
95434	SCASI	228003	SCID due to CORO1A deficiency	295197	SD2b
85297	SCAX3		295199	SD2c	
85292	SCAX4	228003	SCID due to coronin-1A deficiency	93404	SD3
314978	SCAX5	420573	SCID due to CTPS1 deficiency	93406	SD5
284400	SCCB	275	SCID due to DCLRE1C deficiency	84064	SD/THE
98967	SCCD	317425	SCID due to DNA-PKcs deficiency	263463	SDCD, CHST3 type
370396	SCCO	397787	SCID due to IKK2 deficiency	168577	sdCHC
98967	SCD	280142	SCID due to LCK deficiency	300869	SDRPL
420402	SCD syndrome	280142	SCID due to lymphocyte-specific protein tyrosine kinase deficiency	811	SDS
449280	Scedosporiosis			373	SDYS
398069	Schaaf-Yang syndrome	33355	SCID with leukopenia	158029	Sea-blue histiocytosis
1383	Schaap-Taylor-Baraitser syndrome	275	SCID, Athabaskan type	1778	Seaver-Cassidy syndrome
71212	SCHAD deficiency	275	SCID, Athabaskan type		
370039	Schauder syndrome	276	SCIDX1	370052	Sebaceous nevus-central nervous system malformations-aplasia cutis congenital-limbal dermoid-pigmented nevus syndrome
93474	Scheie syndrome	185	Scimitar syndrome		
2353	Schilbach-Rott syndrome	70573	SCLC	370052	Sebaceous nevus-CNS malformations-aplasia cutis congenital-limbal dermoid-pigmented nevus syndrome
59298	Schilder disease	352763	Scleredema		
59298	Schilder's disease	75840	Scleroatonic muscular dystrophy	370052	Sebaceous nevus-CNS malformations-aplasia cutis congenital-limbal dermoid-pigmented nevus syndrome
1830	Schimke immuno-osseous dysplasia	384	Scleroatrophic syndrome		
1830	Schimke syndrome	167635	Scleromyxedema	→182050	Sebastian syndrome
2612	Schimmelpenning syndrome	90400	Scleromyxedema without monoclonal gammopathy		
3137	Schindler disease	75325	Sclerosing dysplasia of bone-ichthyosis-premature ovarian failure syndrome	841	Sebocystomatosis
79279	Schindler disease type 1			168606	Seborrhea-like dermatitis with psoriasiform elements
79280	Schindler disease type 2			79480	Seborrheic pemphigus
79281	Schindler disease type 3			98873	SEC23B-CDG
3138	Schinzel syndrome	63999	Sclerosing mediastinitis	808	Seckel syndrome
798	Schinzel-Giedion syndrome	238593	Sclerosing mesenteritis	141022	Second branchial cleft anomaly
63862	Schisis association	100001	Sclerosing perineurioma		
1247	Schistosomiasis	3152	Scleroosteosis	141022	Second branchial cleft cyst
799	Schizencephaly	384	Sclerolytosis	141022	Second branchial cleft fistula
98973	Schlichting dystrophy	188	SCLS	139420	Secondary acute transverse myelitis
3143	Schmidt syndrome	331176	SCN4	85445	Secondary amyloidosis
2252	Schmitt-Gillenwater-Kelly syndrome	439746	Scondary PAN	169618	Secondary central precocious puberty
3144	Schneckenbecken dysplasia	466677	Scorpion envenomation		
37748	Schnitzler syndrome	832	SCOT deficiency	314962	Secondary HES
98967	Schnyder corneal dystrophy	1514	Scott craniodigital syndrome	314962	Secondary hypereosinophilic syndrome
98967	Schnyder crystalline corneal dystrophy	806	Scott syndrome		
98967	Schnyder crystalline corneal dystrophy	1514	Scott-Bryant-Graham syndrome	2615	Secondary hypertrophic osteoperiostosis with pernio
98967	Schnyder crystalline dystrophy sine crystals	1509	Scott-Taor syndrome		
98967	Schnyder crystalline dystrophy sine crystals	86813	SCRA	140286	Secondary hypoparathyroidism due to impaired parathormon secretion
3145	Schofer-Beetz-Bohl syndrome	83317	Scrub typhus		
3041	Scholte-Begeer-van Essen syndrome	794	SCS	90363	Secondary intestinal lymphangiectasia

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399180	Secondary non-traumatic avascular necrosis	100069	Semantic dementia	3157	Septo-optic dysplasia
399180	Secondary non-traumatic AVN	100069	Semantic primary progressive aphasia	3157	Septo-optic dysplasia spectrum
3452	Secondary non-tropical sprue	100069	Semantic variant PPA	280195	Septopreoptic holoprosencephaly
420259	Secondary PAP	93356	SEMD type 2	280195	Septopreoptic HPE
439746	Secondary periarteritis nodosa	171866	SEMD, aggrecan type	139466	SERKAL syndrome
439746	Secondary polyarteritis nodosa	93351	SEMD, Irapa type	43116	Serotonergic syndrome
420259	Secondary pulmonary alveolar proteinosis	156728	SEMD, MATN3-related	43116	Serotonin storm
99930	Secondary pulmonary hemosiderosis	156728	SEMD, matrilin-3 type	43116	Serotonin syndrome
447774	Secondary sclerosing cholangitis	93356	SEMD, Missouri type	43116	Serotonin toxicity
95427	Secondary short bowel syndrome	93352	SEMD, Shohat type	43116	Serotonin toxidrome
99857	Secondary syringomyelia	93359	SEMD-JL	424073	Serous cystadenocarcinoma of pancreas
364055	SECORD	93360	SEMD-MD	206470	Serous or mucinous cystadenoma of childhood
459051	SED, Stanescu type	93359	SEMDJL1	168829	Serous surface papillary carcinoma
163654	SED-BDS	93360	SEMDJL2	→955	Serpentine fibula-polycystic kidneys syndrome
94068	SEDC	420402	Semicircular canal dehiscence syndrome	35686	Serpiginous choroiditis
567	Sedlackova syndrome	220386	Semilobar holoprosencephaly	157798	Serrated polyposis
647	Seemanova syndrome type 2	842	Seminoma of testis	75508	Servelle-Martorell syndrome
2528	Seemanova-Lesny syndrome	842	Seminomatous germ cell tumor of testis	199343	SeSAME syndrome
251618	SEGA	329284	SENA	1807	Setleis syndrome
2759	Seghers syndrome	79480	Senear-Usher syndrome	85165	Severe achondroplasia-developmental delay-acanthosis nigricans syndrome
67039	Segmental odontomaxillary dysplasia	397596	Senescent T-cells-lymphadenopathy-immunodeficiency syndrome due to p110delta-activating mutation	438207	Severe autosomal recessive macrothrombocytopenia
137608	Segmental outgrowth-lipomatosis-arteriovenous malformation-epidermal nevus syndrome	1369	Sengers syndrome	314911	Severe Canavan disease
314662	Segmental progressive overgrowth syndrome with fibroadipose hyperplasia	2183	Sengers-Hamel-Otten syndrome	277	Severe combined immunodeficiency due to adenosine deaminase deficiency
455	SEI	330001	Senile systemic amyloidosis	275	Severe combined immunodeficiency due to ARTEMIS deficiency
35069	Seitelberger disease	1292	Senior syndrome	357237	Severe combined immunodeficiency due to CARD11 deficiency
79156	Seizures-intellectual disability due to hydroxylysinuria syndrome	84081	Senior-Boichis syndrome	331206	Severe combined immunodeficiency due to complete RAG1/2 deficiency
466926	Seizures-scoliosis-macrocephaly syndrome	3156	Senior-Loken syndrome	228003	Severe combined immunodeficiency due to CORO1A deficiency
199343	Seizures-sensorineural deafness-ataxia-intellectual disability-electrolyte imbalance syndrome	1515	Sensenbrenner syndrome	228003	Severe combined immunodeficiency due to coronin-1A deficiency
35858	Selective cobalamin malabsorption with proteinuria	217622	Sensorineural deafness with dilated cardiomyopathy	420573	Severe combined immunodeficiency due to CTPS1 deficiency
183675	Selective IgG subclass deficiency	857	Sensorineural deafness with imperforate anus and hypoplastic thumbs	275	Severe combined immunodeficiency due to DCLRE1C deficiency
331235	Selective IgM deficiency	217622	Sensorineural hearing loss with dilated cardiomyopathy	317425	Severe combined immunodeficiency due to DNA-PKcs deficiency
331235	Selective immunoglobulin M deficiency	66633	Sensorineural hearing loss-early graying-essential tremor syndrome	397787	Severe combined immunodeficiency due to IKK2 deficiency
165994	Selective pituitary resistance to thyroid hormone	97229	Sensorineural hearing loss-pontobulbar palsy syndrome	280142	Severe combined immunodeficiency due to LCK deficiency
99798	Selective tooth agenesis	70595	Sensory ataxic neuropathy-dysarthria-ophthalmoparesis syndrome	280142	Severe combined immunodeficiency due to lymphocyte-specific protein tyrosine kinase deficiency
281122	Self-healing collodion baby	477	Senter syndrome		
90397	Self-healing papular mucinosis	90118	SEOAN due to MFN2 deficiency		
65748	Self-healing squamous epithelioma type 1	70594	Sepiapterin reductase deficiency		
→1768	Selig-Benacerraf-Greene syndrome	90051	Sepsis in premature infants		
3232	Sellars-Beighton syndrome	180154	Septate vagina		
		137839	Septic phlebitis of the internal jugular vein		

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33355	Severe combined immunodeficiency with leukopenia	169802	Severe hemophilia A	33069	Severe myoclonus epilepsy of infancy
		169793	Severe hemophilia B		
275	Severe combined immunodeficiency, Athabaskan type	467176	Severe hypotonia-psychomotor developmental delay-strabismus-cardiac septal defect syndrome	314655	Severe neonatal hypotonia-seizures-encephalopathy syndrome due to 5q31.3 microdeletion
275	Severe combined immunodeficiency, Athabaskan type	98920	Severe infantile axonal neuropathy with respiratory failure type 1	397593	Severe neonatal lactic acidosis due to NFS1-ISD11 complex deficiency
209370	Severe congenital encephalopathy due to MECP2 mutation	404521	Severe infantile axonal neuropathy with respiratory failure type 2	209370	Severe neonatal-onset encephalopathy with microcephaly
300298	Severe congenital hypochromic anemia with ringed sideroblasts	280763	Severe intellectual disability and progressive spastic paraplegia	363400	Severe neurodegenerative syndrome due to BSLC2 deficiency
300298	Severe congenital hypochromic sideroblastic anemia	420561	Severe intellectual disability-aplasia/hypoplasia of thumb and hallux syndrome	363400	Severe neurodegenerative syndrome with lipodystrophy
171430	Severe congenital nemaline myopathy			216812	Severe osteogenesis imperfecta
99749	Severe congenital neutropenia type 3	466688	Severe intellectual disability-corpora callosa agenesis-facial dysmorphism-cerebellar ataxia syndrome	411543	Severe phosphoribosylpyrophosphate synthetase superactivity
331176	Severe congenital neutropenia type 4			280210	Severe PMD
331176	Severe congenital neutropenia-pulmonary hypertension-superficial venous angiectasis syndrome	94066	Severe intellectual disability-epilepsy-anal anomalies-distal phalangeal hypoplasia	468726	Severe primary trimethylaminuria
369992	Severe dermatitis-multiple allergies-metabolic wasting syndrome	438178	Severe intellectual disability-epilepsy-cataract syndrome due to FAR1 deficiency	411543	Severe PRPP synthetase superactivity
→300751	Severe dilated cardiomyopathy due to lamin A/C mutation			411543	Severe PRPS1 superactivity
→300751	Severe dilated cardiomyopathy with or without myopathy	438178	Severe intellectual disability-epilepsy-cataract syndrome due to fatty acyl-CoA reductase 1 deficiency	163703	Severe refractory status epilepticus owing to presumed encephalitis
98896	Severe dystrophinopathy, Duchenne type	438178	Severe intellectual disability-epilepsy-cataract syndrome due to peroxisomal disorder	169095	Severe T-cell immunodeficiency-congenital alopecia-nail dystrophy syndrome
364055	Severe early-childhood-onset retinal dystrophy			3078	Severe X-linked intellectual disability, Gustavson type
228374	Severe early-onset axonal neuropathy due to light neurofilament subunit deficiency	436141	Severe intellectual disability-hypotonia-strabismus-coarse face-planovalgus syndrome	238329	Severe X-linked mitochondrial encephalomyopathy
90118	Severe early-onset axonal neuropathy due to MFN2 deficiency	363686	Severe intellectual disability-poor language-strabismus-grimacing face-long fingers syndrome	363489	Sex cord-stromal tumor of testis
228374	Severe early-onset axonal neuropathy due to NEFL deficiency	397933	Severe intellectual disability-progressive postnatal microcephaly-midline stereotypic hand movements syndrome	139466	Sex reversion-kidneys, adrenal and lung dysgenesis syndrome
329249	Severe early-onset obesity-insulin resistance syndrome due to SH2B1 deficiency			3162	Sézary lymphoma
440427	Severe early-onset pulmonary alveolar proteinosis due to MARS deficiency	404473	Severe intellectual disability-progressive spastic diplegia syndrome	3162	Sézary syndrome
169793	Severe factor IX deficiency	391307	Severe intellectual disability-short stature-behavioral abnormalities-facial dysmorphism syndrome	373	SGBS
169802	Severe factor VIII deficiency			373	SGBS1
352577	Severe feeding difficulties-failure to thrive-microcephaly due to ASXL3 deficiency syndrome	324307	Severe lateral tibial bowing with short stature	79022	SGBS2
79408	Severe generalized RDEB	2879	Severe limb deficit	457083	SGF
79408	Severe generalized recessive dystrophic epidermolysis bullosa	1236	Severe microbrachycephaly-intellectual disability-athetoid cerebral palsy syndrome	2063	SGFLD syndrome
2109	Severe Hallermann-Streif-François syndrome	369939	Severe motor and intellectual disabilities-sensorineural deafness-dystonia syndrome	35710	SGLT1 deficiency
466026	Severe hemolytic anemia due to G6PD deficiency			69076	SGLT2 deficiency
		33069	Severe myoclonic epilepsy of infancy	2462	SGS
				798	SGS
				2407	Shabbir syndrome
				897	Shah-Waardenburg syndrome
				29822	Shapiro syndrome
				1506	Sharma-Kapoor-Ramji syndrome
				809	Sharp syndrome
				281122	SHCB
				91355	Sheehan syndrome
				1147	Sheldon-Hall syndrome
				3329	SHFLD syndrome
				2440	SHFM

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3329	SHFM associated with aplasia of long bones	435804	Short stature-advanced bone age-early-onset osteoarthritis syndrome	294998	Short toes
90038	Shiga-like toxin-associated HUS	397623	Short stature-auditory canal atresia-mandibular hypoplasia-skeletal anomalies syndrome	295134	Short toes, bilateral
810	Shigellosis			295132	Short toes, unilateral
158014	SHML			357175	Short ulna-dysmorphism-hypotonia-intellectual disability syndrome
1008	Shokeir syndrome	464288	Short stature-brachydactyly-obesity-global developmental delay syndrome	57145	Short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing
99063	Shone complex			935	Short-limb skeletal dysplasia with severe combined immunodeficiency
251515	Short Achilles tendon			79157	Short/branched-chain acyl-coA dehydrogenase deficiency
26792	Short chain acyl-CoA dehydrogenase deficiency	2994	Short stature-craniofacial anomalies-genital hypoplasia syndrome	2580	Shoulder and girdle defects-familial intellectual disability syndrome
66518	Short fifth metacarpals-insulin resistance syndrome	2866	Short stature-deafness-neutrophil dysfunction-dysmorphism syndrome	1940	Shoulder and thorax deformity-congenital heart disease syndrome
294996	Short fingers	171706	Short stature-delayed bone age due to thyroid hormone metabolism deficiency	314795	SHOX-related short stature
295130	Short fingers, bilateral			567	Shprintzen syndrome
295128	Short fingers, unilateral			2462	Shprintzen-Goldberg syndrome
935	Short limb skeletal dysplasia with SCID	2332	Short stature-facial and skeletal anomalies-intellectual disability-macrodontia syndrome	3165	Shulman syndrome
93270	Short rib-polydactyly syndrome type 1			811	Shwachman syndrome
93269	Short rib-polydactyly syndrome type 2			811	Shwachman-Bodian-Diamond syndrome
93271	Short rib-polydactyly syndrome type 3	→2995	Short stature-intellectual disability-eye anomalies-cleft lip/palate syndrome	811	Shwachman-Diamond syndrome
93268	Short rib-polydactyly syndrome type 4	420794	Short stature-kyphosis-hypoplasia of basal ilia-cone epiphyses-facial dysmorphism syndrome	812	Sialidosis type 1
93268	Short rib-polydactyly syndrome, Beemer-Langer type			87876	Sialidosis type 2
93269	Short rib-polydactyly syndrome, Majewski type			3166	Sialuria
93270	Short rib-polydactyly syndrome, Saldino-Noonan type	1937	Short stature-locking fingers syndrome	3166	Sialuria, French type
93271	Short rib-polydactyly syndrome, Verma-Naumoff type	423454	Short stature-nail dysplasia-marginal palmoplantar keratoderma-oral hyperpigmentation syndrome	98920	SIANRF
→1263	Short ribs-craniosynostosis-polysyndactyly syndrome			→33364	SIBIDS syndrome
314811	Short stature due to GHSR deficiency			611	SIBM
629	Short stature due to growth hormone qualitative anomaly	3102	Short stature-Pierre Robin sequence-cleft mandible-hand anomalies clubfoot syndrome	232	Sickle cell anemia
633	Short stature due to growth hormone resistance			232	Sickle cell disease
314811	Short stature due to growth hormone secretagogue receptor deficiency			251359	Sickle cell-beta-thalassemia disease syndrome
632	Short stature due to isolated growth hormone deficiency with X-linked hypogammaglobulinemia	391677	Short stature-optic atrophy-Pelger-Huët anomaly syndrome	251365	Sickle cell-hemoglobin C disease syndrome
314802	Short stature due to partial GHR deficiency	2868	Short stature-valvular heart disease-characteristic facies syndrome	251370	Sickle cell-hemoglobin D disease syndrome
314802	Short stature due to partial growth hormone receptor deficiency			251375	Sickle cell-hemoglobin E disease syndrome
140941	Short stature due to primary acid-labile subunit deficiency			210272	Sickness of disembarkment
220465	Short stature due to STAT5b deficiency	85442	Short stature-pituitary and cerebellar defects-small sella turcica syndrome	838	SICRET syndrome
2867	Short stature, Brussels type	2868	Short stature-wormian bones-dextrocardia syndrome	168593	SIDDT
		2865	Short stature-webbed neck-heart disease syndrome	54028	Sideropenic dysphagia
		2863	Short stature-wormian bones-dextrocardia syndrome	2267	Sidransky-Feinstein-Goodman syndrome
		3163	SHORT syndrome	3167	Siegler-Brewer-Carey syndrome
		2832	Short tarsus-absence of lower eyelashes syndrome	→244	Siewert syndrome
		251515	Short tendo calcaneus	369861	SIFD syndrome
				314786	Silent pituitary adenoma
				71276	Silent sinus syndrome
				3168	Sillence syndrome
				60014	Silver staining
				100998	Silver syndrome

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
813	Silver-Russell dwarfism	800	SJS	363447	SMALED
813	Silver-Russell syndrome	800	SJS1	209341	SMALED1
231137	Silver-Russell syndrome due to 7p11.2-p13 microduplication	3206	SJS2	363454	SMALED2
		95455	SJS-TEN	284400	Small cell bladder cancer
231137	Silver-Russell syndrome due to 7p11.2p13 microduplication	1426	Skeletal dysplasia, Greenberg type	284400	Small cell bladder carcinoma
		2565	Skeletal dysplasia-brachydactyly syndrome	284400	Small cell carcinoma of the bladder
231144	Silver-Russell syndrome due to 11p15 microduplication	1858	Skeletal dysplasia-epilepsy-short stature syndrome	370396	Small cell carcinoma of the ovary
397590	Silver-Russell syndrome due to a point mutation			284400	Small cell carcinoma of the urinary bladder
231140	Silver-Russell syndrome due to an imprinting defect of 11p15	1436	Skeletal dysplasia-intellectual disability syndrome	70573	Small cell lung cancer
				370396	Small cell ovarian carcinoma
231137	Silver-Russell syndrome due to dup(7)(p11.2p13)	477831	Skeletal overgrowth-craniofacial dysmorphism-hyperelastic skin-white matter lesions syndrome	838	Small infarctions of cochlear, retinal and encephalic tissue
96182	Silver-Russell syndrome due to maternal uniparental disomy of chromosome 7			1201	Small intestinal atresia
		293165	Skin fragility-woolly hair-palmoplantar hyperkeratosis syndrome	67038	Small lymphocytic lymphoma
231147	Silver-Russell syndrome due to maternal uniparental disomy of chromosome 11	293165	Skin fragility-woolly hair-palmoplantar keratoderma syndrome	543	Small non-cleaved cell lymphoma
				1509	Small patella syndrome
231137	Silver-Russell syndrome due to trisomy 7p11.2-p13	178475	Skin infectious botulism	466962	SMARCA4-deficient sarcoma of thorax
		178475	Skin toxin-mediated botulism	466962	SMARCA4-deficient thoracic sarcoma
231137	Silver-Russell syndrome due to trisomy 7p11.2p13	52503	SLC6A8 deficiency	98920	SMARD1
		238459	SLC35A1-CDG	404521	SMARD2
1968	Simosa-Penhaszadeh-Bustos syndrome	356961	SLC35A2-CDG	481	SMA1
		370943	SLC35A3-CDG	1145	SMA2
91139	Simple cryoglobulinemia	99843	SLC35C1-CDG	139557	SMA3
373	Simpson dysmorphia syndrome	3144	SLC35D1-CDG	98959	SMCD
373	Simpson-Golabi-Behmel syndrome	468699	SLC39A8 deficiency	85167	SMD-CRD
373	Simpson-Golabi-Behmel syndrome type 1	468699	SLC39A8-CDG	33069	SMEI
		93552	SLE, pediatric onset	93974	Smith-Fineman-Myers syndrome
79022	Simpson-Golabi-Behmel syndrome type 2	3385	Sleeping sickness	818	Smith-Lemli-Opitz syndrome
97337	Sinding-Larsen-Johansson disease	88633	SLK	819	Smith-Magenis syndrome
		818	SLOS	178355	Smith-McCort dysplasia
50809	Singh-Williams-McAlister syndrome	70472	SLSJ-COX deficiency	2286	SMMCI
2286	Single upper central incisor	3156	SLSN	158775	Smouldering systemic mastocytosis
439755	Single-organ PAN	584	Sly disease	3198	SMS
439755	Single-organ periarteritis nodosa	70	SMA	86854	SMZL
439755	Single-organ polyarteritis nodosa	83330	SMA1	449285	Snakebite envenomation
85191	Singleton-Merten dysplasia	83418	SMA2	820	Sneddon syndrome
85191	Singleton-Merten syndrome	83419	SMA3	48377	Sneddon-Wilkinson disease
1260	Sino-auricular heart block	83420	SMA4	91496	Snowflake vitreoretinal degeneration
324321	Sinoatrial node dysfunction and deafness	83330	SMA type 1	3063	Snyder-Robinson syndrome
		83418	SMA type 2	3157	SOD
158014	Sinus histiocytosis with massive lymphadenopathy	83419	SMA type 3	67039	SOD
		83420	SMA type 4	306577	Sodium channelopathy-related small fiber neuropathy
890	Sinusoidal obstruction syndrome	83330	SMA type I	99903	Sudoku
247698	Sipple syndrome	83418	SMA type II	314394	SOFT syndrome
3169	Sirenomelia	83419	SMA type III	100002	Soft tissue perineurioma
2882	Sitosterolemia	83420	SMA type IV	2234	Sohval-Soffer syndrome
157769	Situs ambiguous	83330	SMA-I	137608	SOLAMEN syndrome
157769	Situs ambiguus	83418	SMA-II	97230	Solar urticaria
101063	Situs inversus	83419	SMA-III		
101063	Situs inversus totalis	83419	SMA-III		
816	Sjögren-Larsson syndrome	83420	SMA-IV		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
424065	Solid pseudopapillary carcinoma of pancreas	2816	Spastic paraplegia-epilepsy-intellectual disability syndrome	352403	Spectrin-associated autosomal recessive cerebellar ataxia
83468	Solitary bone cyst	2819	Spastic paraplegia-facial-cutaneous lesions syndrome	352403	Spectrin-associated autosomal recessive cerebellar ataxia type 1
2126	Solitary fibrous tumor				
79455	Solitary mastocytoma	2818	Spastic paraplegia-glaucoma-intellectual disability syndrome	209908	Speech and language disorder with orofacial dyspraxia
2286	Solitary median maxillary central incisor syndrome	2822	Spastic paraplegia-intellectual disability-thin corpus callosum syndrome	209908	Speech-language disorder type 1
100035	Solitary necrotic tumor of the liver			→2909	Spellacy-Gibbs-Watts syndrome
86855	Solitary plasmacytoma	2820	Spastic paraplegia-nephritis-deafness syndrome	2816	SPEMR
209964	Solitary rectal ulcer syndrome			1855	SPENCD
2612	Solomon syndrome	2821	Spastic paraplegia-neuropathy-poikiloderma syndrome	→1855	SPENCDI
314769	Somatomammotropinoma			99865	Spermatocytic seminoma
97283	Somatostatinoma	320406	Spastic paraplegia-optic atrophy-neuropathy syndrome	306617	SPG1
2564	Sommer-Hines syndrome			99015	SPG2
1064	Sommer-Rathbun-Battles syndrome	329475	Spastic paraplegia-Paget disease of bone syndrome	100985	SPG4
1529	Sommer-Young-Wee-Frye syndrome			100986	SPG5A
1355	Sonoda syndrome	2826	Spastic paraplegia-precocious puberty syndrome	100988	SPG6
391677	SOPH syndrome			99013	SPG7
1471	Sorsby syndrome	464282	Spastic paraplegia-psychomotor retardation-seizures syndrome	100989	SPG8
59181	Sorsby's fundus dystrophy			100990	SPG9
821	Sotos syndrome	100996	Spastic paraplegia-retinal degeneration syndrome	100991	SPG10
420179	Sotos syndrome 2			2822	SPG11
98868	Southeast Asian ovalocytosis	464282	Spastic paraplegia-severe developmental delay-epilepsy syndrome	100993	SPG12
352403	SPARCA			100994	SPG13
352403	SPARCA1	3011	Spastic quadriplegia-retinitis pigmentosa-intellectual disability syndrome	100995	SPG14
79132	Sparse hair-short stature-skin anomalies syndrome			100996	SPG15
279882	Spasmus nutans	447997	Spastic quadriplegia-thin corpus callosum-progressive postnatal microcephaly syndrome	100997	SPG16
1182	Spastic ataxia with congenital miosis			100998	SPG17
2572	Spastic ataxia-corneal dystrophy syndrome	210141	Spastic quadriplegic cerebral palsy	209951	SPG18
2572	Spastic ataxia-ocular anomalies syndrome	3011	Spastic tetraplegia-retinitis pigmentosa-intellectual disability syndrome	100999	SPG19
99015	Spastic gait type 2			101000	SPG20
99015	Spastic paraparesis type 2	447997	Spastic tetraplegia-thin corpus callosum-progressive postnatal microcephaly syndrome	101001	SPG21
100990	Spastic paraparesis-amyopathy-cataracts-gastroesophageal reflux syndrome			101003	SPG23
2815	Spastic paraparesis-deafness syndrome	401866	Spasticity-ataxia-gait anomalies syndrome	101004	SPG24
101003	Spastic paraparesis-vitiligo-premature graying-characteristic facies syndrome	251282	SPAX1	101005	SPG25
		397946	SPAX2	101006	SPG26
		314603	SPAX3	101007	SPG27
		254343	SPAX4	101008	SPG28
139480	Spastic paraplegia due to neuropathy target esterase mutation	313772	SPAX5	101009	SPG29
		98	SPAX6	101010	SPG30
		1182	SPAX7	101011	SPG31
139480	Spastic paraplegia due to NTE mutation	158	SPCD	171622	SPG32
		295195	SPD1	171607	SPG34
431329	Spastic paraplegia due to partial TFG deficiency	295197	SPD2	171629	SPG35
		295199	SPD3	320365	SPG36
99015	Spastic paraplegia type 2	98	SPAX6	171612	SPG37
99013	Spastic paraplegia type 7	1182	SPAX7	171617	SPG38
100998	Spastic paraplegia-amyotrophy of hands and feet	158	SPCD	139480	SPG39
		295195	SPD, Vordingborg type	295195	SPD1
		295197	SPD2	320355	SPG41
		295199	SPD3	171863	SPG42
		295197	SPD, Debeer type	320370	SPG43
		295199	SPD, Malik type	320401	SPG44
		295195	SPD, Vordingborg type	320396	SPG45
				320391	SPG46

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
306511	SPG48	1955	Spinocerebellar ataxia and erythrokeratodermia	64753	Spinocerebellar ataxia with axonal neuropathy type 2
320385	SPG49			254881	Spinocerebellar ataxia with epilepsy
319199	SPG53	412057	Spinocerebellar ataxia autosomal recessive type 16	276241	Spinocerebellar ataxia, Thomas type
320380	SPG54			2074	Spinocerebellar ataxia-amyotrophy-deafness syndrome
320375	SPG55	453521	Spinocerebellar ataxia autosomal recessive type 17	1185	Spinocerebellar ataxia-dysmorphism syndrome
320411	SPG56				
431329	SPG57	404493	Spinocerebellar ataxia autosomal recessive type 23	3177	Spinocerebellar degeneration-corneal dystrophy syndrome
397946	SPG58				
401795	SPG59	98755	Spinocerebellar ataxia type 1	99903	Spirillary rat-bite fever
401800	SPG60	94124	Spinocerebellar ataxia type 1 with axonal neuropathy		
401780	SPG61	98756	Spinocerebellar ataxia type 2	757	Spitzer-Weinstein syndrome
401785	SPG62	98757	Spinocerebellar ataxia type 3	300869	Splenic diffuse red pulp lymphoma
401805	SPG63	276238	Spinocerebellar ataxia type 3, Joseph type	300869	Splenic diffuse red pulp small B-cell lymphoma
401810	SPG64				
320396	SPG65	276244	Spinocerebellar ataxia type 3, Machado type	86854	Splenic marginal zone lymphoma
401815	SPG66				
401820	SPG67	98765	Spinocerebellar ataxia type 4	2063	Splenogonadal fusion-limb defects-micrognathia syndrome
401825	SPG68	98766	Spinocerebellar ataxia type 5	47612	Splenomegaly-neutropenia-rheumatoid arthritis syndrome
401830	SPG69				
401835	SPG70	98758	Spinocerebellar ataxia type 6	294994	Split foot
401840	SPG71	94147	Spinocerebellar ataxia type 7	2439	Split foot deformity-mandibulofacial dysostosis syndrome
401849	SPG72	98760	Spinocerebellar ataxia type 8		
444099	SPG73	98761	Spinocerebellar ataxia type 10	295126	Split foot, bilateral
468661	SPG74	98767	Spinocerebellar ataxia type 11	295124	Split foot, unilateral
459056	SPG75	98762	Spinocerebellar ataxia type 12	294992	Split hand
466722	SPG77	98768	Spinocerebellar ataxia type 13	2440	Split hand foot malformation
268129	Spheroid body myopathy	98763	Spinocerebellar ataxia type 14	2437	Split hand with obstructive uropathy, spina bifida and diaphragmatic defects
3449	Spherophakia-brachymorphia syndrome	98769	Spinocerebellar ataxia type 15/16		
306553	Spherulocytosis	→98769	Spinocerebellar ataxia type 16	295122	Split hand, bilateral
79264	Spielmeier-Vogt disease	98759	Spinocerebellar ataxia type 17	295120	Split hand, unilateral
314432	Spigelian hernia-cryptorchidism syndrome	98771	Spinocerebellar ataxia type 18	2440	Split hand-split foot malformation
268369	Spina bifida aperta	98772	Spinocerebellar ataxia type 19/22	71271	Split hand-split foot-deafness syndrome
3176	Spina bifida-hypospadias syndrome	101110	Spinocerebellar ataxia type 20		
53721	Spinal arteriovenous metamerism syndrome	98773	Spinocerebellar ataxia type 21	2437	Split hand-urinary anomalies-spina bifida syndrome
1217	Spinal atrophy-ophthalmoplegia-pyramidal syndrome	→98772	Spinocerebellar ataxia type 22	3329	Split hand/foot malformation with long bone deficiency
90058	Spinal cord injury	101108	Spinocerebellar ataxia type 23		
1145	Spinal muscular atrophy with arthrogyposis	101111	Spinocerebellar ataxia type 25	958	Split hand/split foot-mandibular hypoplasia syndrome
98920	Spinal muscular atrophy with respiratory distress type 1	101112	Spinocerebellar ataxia type 26	2329	Split hand/split foot-nystagmus syndrome
404521	Spinal muscular atrophy with respiratory distress type 2	98764	Spinocerebellar ataxia type 27		
83420	Spinal muscular atrophy, adult form	101109	Spinocerebellar ataxia type 28	1756	Split notochord syndrome
73245	Spinal muscular atrophy-Dandy-Walker malformation-cataracts syndrome	208513	Spinocerebellar ataxia type 29	3329	Split-hand/foot malformation associated with aplasia of long bones
2590	Spinal muscular atrophy-progressive myoclonic epilepsy syndrome	211017	Spinocerebellar ataxia type 30		
210584	Spindle cell hemangioma	217012	Spinocerebellar ataxia type 31	320406	SPOAN
210584	Spindle cell hemangioma	276183	Spinocerebellar ataxia type 32		
		1955	Spinocerebellar ataxia type 34	93357	SPONASTRIME dysplasia
		276193	Spinocerebellar ataxia type 35		
		276198	Spinocerebellar ataxia type 36	1190	Spondylo-humero-femoral dysplasia
		363710	Spinocerebellar ataxia type 37	228387	Spondylo-megaepiphyseal-metaphyseal dysplasia
		423296	Spinocerebellar ataxia type 38		
		423275	Spinocerebellar ataxia type 40	85194	Spondylo-ocular syndrome
		458798	Spinocerebellar ataxia type 41		
		458803	Spinocerebellar ataxia type 42	3180	Spondylocamptodactyly syndrome
		363710	Spinocerebellar ataxia with altered vertical eye movements		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
3275	Spondylorcarpotarsal synostosis	93357	Spondyloepimetaphyseal dysplasia, Sponastrime type	→1855	Spondylometaphyseal dysplasia with combined immunodeficiency
94095	Spondylocostal dysostosis-anal and genitourinary malformations syndrome	168451	Spondyloepimetaphyseal dysplasia-abnormal dentition syndrome	1855	Spondylometaphyseal dysplasia with enchondromatous changes
329252	Spondylocostal dysostosis-hypospadias-intellectual disability syndrome	168443	Spondyloepimetaphyseal dysplasia-hypotrichosis syndrome	93316	Spondylometaphyseal dysplasia with severe genu valgum
1855	Spondyloenchondrodysplasia	93358	Spondyloepimetaphyseal dysplasia-short limb-abnormal calcification syndrome	168555	Spondylometaphyseal dysplasia, A4 type
→1855	Spondyloenchondrodysplasia with immune dysregulation	94068	Spondyloepiphyseal dysplasia congenita	93316	Spondylometaphyseal dysplasia, Algerian type
1855	Spondyloenchondromatosis	93284	Spondyloepiphyseal dysplasia tarda	93315	Spondylometaphyseal dysplasia, 'corner fracture' type
93346	Spondyloepimetaphyseal dysplasia congenita, Strudwick type	163665	Spondyloepiphyseal dysplasia tarda, Kohn type	370019	Spondylometaphyseal dysplasia, Czarny-Ratajczak type
93356	Spondyloepimetaphyseal dysplasia type 2	1159	Spondyloepiphyseal dysplasia tarda-progressive arthropathy syndrome	168544	Spondylometaphyseal dysplasia, Golden type
93360	Spondyloepimetaphyseal dysplasia with joint laxicity, Hall type	263463	Spondyloepiphyseal dysplasia with congenital joint dyslocations, CHST3 type	93314	Spondylometaphyseal dysplasia, Kozlowski type
93359	Spondyloepimetaphyseal dysplasia with joint laxity	→93284	Spondyloepiphyseal dysplasia, Byers type	93316	Spondylometaphyseal dysplasia, Schmidt type
93359	Spondyloepimetaphyseal dysplasia with joint laxity type 1	163654	Spondyloepiphyseal dysplasia, Cantu type	93317	Spondylometaphyseal dysplasia, Sedaghatian type
93360	Spondyloepimetaphyseal dysplasia with joint laxity type 2	93283	Spondyloepiphyseal dysplasia, Kimberley type	93315	Spondylometaphyseal dysplasia, Sutcliffe type
93360	Spondyloepimetaphyseal dysplasia with joint laxity, leptodactylic type	163668	Spondyloepiphyseal dysplasia, MacDermot type	168552	Spondylometaphyseal dysplasia-bowed forearms-facial dysmorphism syndrome
93360	Spondyloepimetaphyseal dysplasia with multiple dislocations	263482	Spondyloepiphyseal dysplasia, Maroteaux type	85167	Spondylometaphyseal dysplasia-cone-rod dystrophy syndrome
171866	Spondyloepimetaphyseal dysplasia, aggrecan type	163649	Spondyloepiphyseal dysplasia, Nishimura type	1856	Spondyloperipheral dysplasia-short ulna syndrome
93347	Spondyloepimetaphyseal dysplasia, anauxetic type	→263463	Spondyloepiphyseal dysplasia, Omani type	141	Spongy degeneration of the brain
168448	Spondyloepimetaphyseal dysplasia, Bieganski type	163662	Spondyloepiphyseal dysplasia, Reardon type	54260	Spongy myocardium
168454	Spondyloepimetaphyseal dysplasia, Geneviève type	459051	Spondyloepiphyseal dysplasia, Stanescu type	443180	Spontaneous cerebrospinal fluid leak
99642	Spondyloepimetaphyseal dysplasia, Handigodu type	163654	Spondyloepiphyseal dysplasia-brachydactyly-speech disorder syndrome	443180	Spontaneous intracranial hypotension
93351	Spondyloepimetaphyseal dysplasia, Irapa type	163649	Spondyloepiphyseal dysplasia-craniosynostosis-cleft palate-cataract-intellectual disability syndrome	29822	Spontaneous periodic hypothermia
370015	Spondyloepimetaphyseal dysplasia, Isidor type	163668	Spondyloepiphyseal dysplasia-myopia-sensorineural deafness syndrome	247234	Sporadic adult-onset ataxia of unknown etiology
156728	Spondyloepimetaphyseal dysplasia, matrilin-3 type	1830	Spondyloepiphyseal dysplasia-nephrotic syndrome	204	Sporadic CJD
93347	Spondyloepimetaphyseal dysplasia, Menger type	→93284	Spondyloepiphyseal dysplasia-punctate corneal dystrophy syndrome	204	Sporadic Creutzfeldt-Jakob disease
93356	Spondyloepimetaphyseal dysplasia, Missouri type	353298	Spondyloepiphyseal dysplasia-retinal dystrophy-immunodeficiency syndrome	1665	Sporadic fetal brain disruption sequence
93282	Spondyloepimetaphyseal dysplasia, Pakistani type			306776	Sporadic hyperekplexia
93282	Spondyloepimetaphyseal dysplasia, PAPSS2 type			225147	Sporadic IBSN
93352	Spondyloepimetaphyseal dysplasia, Shohat type			84271	Sporadic idiopathic nephrosis
				84271	Sporadic idiopathic steroid-resistant nephrotic syndrome
				97555	Sporadic idiopathic steroid-resistant nephrotic syndrome with collapsing glomerulopathy
				93222	Sporadic idiopathic steroid-resistant nephrotic syndrome with diffuse mesangial proliferation

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
93220	Sporadic idiopathic steroid-resistant nephrotic syndrome with diffuse mesangial sclerosis	424039	Squamous cell carcinoma of pancreas	3184	Steatocystoma multiplex-natal teeth syndrome
		398058	Squamous cell carcinoma of penis	438117	Steel syndrome
93218	Sporadic idiopathic steroid-resistant nephrotic syndrome with focal segmental glomerulosclerosis	424002	Squamous cell carcinoma of rectum	240071	Steele-Richardson-Olszewski disease
		418959	Squamous cell carcinoma of stomach	565	Steely hair disease
93218	Sporadic idiopathic steroid-resistant nephrotic syndrome with focal segmental hyalinosis	424019	Squamous cell carcinoma of the anal canal	565	Steely hair syndrome
				273	Steinert disease
93221	Sporadic idiopathic steroid-resistant nephrotic syndrome with minimal changes	213767	Squamous cell carcinoma of the cervix uteri	273	Steinert myotonic dystrophy
		213716	Squamous cell carcinoma of the corpus uteri	3186	Steinfeld syndrome
611	Sporadic inclusion body myositis			168953	Stem cell leukemia/lymphoma
225147	Sporadic infantile bilateral striatal necrosis	457252	Squamous cell carcinoma of the oral tongue	99087	Stenosis or atrophy of the coronary ostium
225147	Sporadic infantile striatonigral degeneration	423968	Squamous cell carcinoma of the small bowel	210115	Sterile multifocal osteomyelitis with periostitis and pustulosis
225147	Sporadic infantile striatonigral necrosis	423968	Squamous cell carcinoma of the small intestine	3194	Stern-Lubinsky-Durrie syndrome
227510	Sporadic olivopontocerebellar atrophy type 1	324737	SRD5A3-CDG	→42775	Sternal malformation-vascular dysplasia syndrome
227510	Sporadic OPCA type 1	83601	SREAT	753	Steroid 5-alpha-reductase deficiency
276624	Sporadic pheochromocytoma	330001	SSA	3196	Steroid dehydrogenase deficiency-dental anomalies syndrome
		22	SSADH deficiency	461	Steroid sulfatase deficiency
276621	Sporadic pheochromocytoma/secretory paraganglioma	466926	SSM syndrome		Steroid-responsive encephalopathy associated with autoimmune thyroiditis
		2806	SSPE	83601	
		50944	SSPS	93207	Steroid-sensitive MCNS
443057	Sporadic porphyria cutanea tarda	370927	SSR4-CDG	→69061	Steroid-sensitive nephrotic syndrome without renal biopsy
276627	Sporadic secreting paraganglioma	2323	SSS	909	Sterol 27-hydroxylase deficiency
826	Sporotrichosis	36236	SSSS	46059	Sterol C5-desaturase deficiency
464282	SPPRS syndrome	83484	St. Louis encephalitis	36426	Stevens-Johnson syndrome
70594	SPR deficiency	2454	Stalker-Chitayat syndrome	828	Stickler syndrome
94068	Spranger-Wiedemann disease	1798	Stanescu osteosclerosis	90653	Stickler syndrome type 1
3181	Sprengel deformity	3235	Stapedo-vestibular ankylosis	90654	Stickler syndrome type 2
70476	Spring catarrh	140917	Stapes ankylosis with broad thumbs and toes	166100	Stickler syndrome type 3
234	Sprinz-Nelson syndrome	36238	Staphylococcal necrotizing pneumonia	166100	Stickler syndrome, non-ocular type
3198	SPS			3197	Stiff baby syndrome
1509	SPS	36236	Staphylococcal scalded skin syndrome	443804	Stiff leg syndrome
431255	SPSMA	36235	Staphylococcal scarlet fever	3198	Stiff man syndrome
86884	SPTCL	99919	Staphylococcal toxic-shock syndrome	3198	Stiff person syndrome and related disorders
51083	SQTS	99919	Staphylococcal TSS	2833	Stiff skin syndrome
423994	Squamous cell carcinoma of colon	140952	STAR syndrome	85414	Still disease
99977	Squamous cell carcinoma of esophagus	827	Stargardt 1	233	Stilling-Turk-Duane syndrome
424996	Squamous cell carcinoma of gallbladder and extrahepatic biliary tract	827	Stargardt disease	3199	Stimmler syndrome
		85146	Stark-Kaesler syndrome	425120	STING-associated vasculopathy with onset in infancy
424996	Squamous cell carcinoma of gallbladder and EBT	166427	Startle epilepsy	2972	Stoeltinga-de Koomen-Davis syndrome
67037	Squamous cell carcinoma of head and neck	391311	STAT1 deficiency	3200	Stoll-Alembik-Finck syndrome
		2314	STAT3 deficiency	3074	Stoll-Géraudel-Chauvin syndrome
424975	Squamous cell carcinoma of liver and IBT	438159	STAT3-related early-onset multisystem autoimmune disease	3201	Stoll-Kieny-Dott syndrome
424975	Squamous cell carcinoma of liver and intrahepatic biliary tract	329284	Static encephalopathy of childhood with neurodegeneration in adulthood	2878	Stoll-Lévy-Francfort syndrome
		841	Steatocystoma multiplex	168577	Stomatatin-deficient cryohydrocytosis
				98868	Stomatocytic elliptocytosis

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
353253	Stomatodynia	356	Subacute spongiform encephalopathy, Gerstmann-Straussler type	308400	Sulfite oxidase deficiency due to molybdenum cofactor deficiency type C
353253	Stomatopyrosis				
337	Stone man syndrome				
3204	Stormorken syndrome	99113	Subaortic course of brachiocephalic vein	99731	Sulfocysteinuria
3204	Stormorken-Sjaastad-Langset syndrome	99113	Subaortic course of innominate vein	65682	Summerskill-Walshe-Tygstrup syndrome
99064	Straddling and/or overriding mitral valve	3191	Subaortic stenosis-short stature syndrome	254395	Summertime actinic lichenoid eruption
95461	Straddling or overriding tricuspid valve	48377	Subcorneal pustular dermatitis	3210	Summitt syndrome
		48377	Subcorneal pustular dermatosis	57145	SUNCT syndrome
1277	Stratton-Garcia-Young syndrome	99796	Subcortical band heterotopia	130	SUNDS
2863	Stratton-Parker syndrome	313808	Subcortical gliosis of Neumann	455	Superficial epidermolytic ichthyosis
99905	Streptobacillary rat-bite fever	99796	Subcortical laminar heterotopia	98961	Superficial granular corneal dystrophy
99918	Streptococcal toxic-shock syndrome	86884	Subcutaneous panniculitic T-cell lymphoma		
99918	Streptococcal TSS			79490	Superficial lymphangioma
66529	Stress cardiomyopathy	86884	Subcutaneous panniculitis-like T-cell lymphoma	79490	Superficial lymphatic malformation
90041	Stress erythrocytosis			247245	Superficial siderosis
90041	Stress polycythemia	251618	Subependymal giant cell astrocytoma	88633	Superior limbic keratoconjunctivitis
50942	Striate palmoplantar keratoderma			155884	Superior palpebral coloboma
137599	Stromal keratitis	101030	Subependymal nodular heterotopia	180182	Supernumerary breasts
213711	Stromal sarcoma of the corpus uteri	251639	Subependymoma	96170	Supernumerary der(22) syndrome
76	Strongyloidiasis	98957	Subepithelial amyloidosis of the cornea	141096	Supernumerary nostril
100984	Strümpell disease	98959	Subepithelial mucinous corneal dystrophy	295002	Supernumerary phalanges
370921	STT3A-CDG			295142	Supernumerary phalanges, bilateral
370924	STT3B-CDG	155878	Submucosal cleft palate	295140	Supernumerary phalanges, unilateral
328	Stuart-Prower factor deficiency			295002	Supernumerary phalanx
3205	Sturge-Weber syndrome	3190	Subpulmonary stenosis	295142	Supernumerary phalanx, bilateral
3205	Sturge-Weber-Dimitri syndrome	1606	Subtelomeric 1p36 deletion	295140	Supernumerary phalanx, unilateral
3205	Sturge-Weber-Krabbe angiomatosis	96168	Subtelomeric deletion 13q34	1450	Supernumerary ring/marker 8
3205	Sturge-Weber-Krabbe syndrome	180129	Subtotal septate uterus	1461	Superoinferior ventricles
3206	Stüve-Wiedemann dysplasia	→2609	Succinic acidemia	764	Suppurative myositis
3206	Stüve-Wiedemann syndrome	22	Succinic semialdehyde dehydrogenase deficiency	466695	Supratip dysplasia
3206	Stüve-Wiedemann/Schwartz-Jampel type 2 syndrome	832	Succinyl-CoA acetoacetate transferase deficiency	3193	Supravalvular aortic stenosis
166277	Suarez-Stickler syndrome	832	Succinyl-CoA:3-ketoacid CoA transferase deficiency	3192	Supravalvular pulmonary stenosis
101029	Sub-cortical nodular heterotopia			391351	SURF1-related Charcot-Marie-Tooth disease type 4
79093	Subacute angiohypertrophic myelomalacia	832	Succinyl-CoA:3-oxoacid CoA transferase deficiency	391351	SURF1-related CMT4
79093	Subacute ascending necrotizing myelitis	702	Sudanophilic leukodystrophy, Paelizeus-Merzbacher type	391351	SURF1-related severe demyelinating Charcot-Marie-Tooth disease
163525	Subacute cutaneous lupus erythematosus	168593	Sudden infant death-dysgenesis of the testes syndrome	838	Susac syndrome
2806	Subacute inclusion body encephalitis			331226	Susceptibility to infection due to TYK2 deficiency
206594	Subacute inflammatory demyelinating polyneuropathy	130	Sudden unexplained nocturnal death syndrome	447740	Susceptibility to localized juvenile periodontitis
206594	Subacute inflammatory demyelinating polyradiculoneuropathy	2752	Sugarman syndrome	169085	Susceptibility to respiratory infections associated with CD8alpha chain mutation
		3412	Sujansky-Leonard syndrome		
98824	Subacute myeloid leukemia	99732	Sulfite oxidase deficiency due to molybdenum cofactor deficiency	391311	Susceptibility to viral and mycobacterial infections
79093	Subacute necrotizing myelitis	308386	Sulfite oxidase deficiency due to molybdenum cofactor deficiency type A	3193	SVAS
2806	Subacute sclerosing leukoencephalitis			86813	Sveinsson chorioretinal atrophy
2806	Subacute sclerosing panencephalitis	308393	Sulfite oxidase deficiency due to molybdenum cofactor deficiency type B	3243	Sweet syndrome
				247165	Swift disease
				247165	Swift-Feer disease

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
3205	SWS	3258	Syndactyly type 7	85274	Syndromic X-linked intellectual disability 7
242	Swyer syndrome	2498	Syndactyly type 8	85279	Syndromic X-linked intellectual disability due to JARID1C mutation
90038	Sxt-HUS	157801	Syndactyly type 9	85295	Syndromic X-linked intellectual disability type 10
306731	Sydenham chorea	157801	Syndactyly, Malik-Percin type	85286	Syndromic X-linked intellectual disability type 11
295138	Symbrachydactyly of hand and foot, bilateral	295012	Syndactyly, mitten type	319332	SYNE1-related AMC
295136	Symbrachydactyly of hand and foot, unilateral	357332	Syndactyly-camptodactyly and clinodactyly of fifth fingers-bifid toes syndrome	319332	SYNE1-related arthrogryposis multiplex congenita
1570	Symbrachydactyly of hands and feet	3253	Syndactyly-ectodermal dysplasia-cleft/lip palate	3262	Syngnathia multiple anomalies
60015	Symmetric parietal foramina	3259	Syndactyly-polydactyly-ear lobe syndrome	3263	Syngnathia-cleft palate syndrome
1314	Symmetrical thalamic calcifications	85203	Syndactyly-preaxial polydactyly-sternal deformity syndrome	35098	Synostotic plagiocephaly
79098	Sympathetic ophthalmia	140952	Syndactyly-telecanthus-anogenital and renal malformations syndrome	3273	Synovial sarcoma
79098	Sympathetic uveitis	→1159	Syndesmodysplasic dwarfism	3273	Synoviosarcoma
3246	Symphalangism with multiple anomalies of hands and feet	2143	Syndromic of ocular and facial anomalies, telecanthus and deafness	793	Synovitis-acne-pustulosis-hyperostosis-osteitis syndrome
3250	Symphalangism, Cushing type	52	Syndromic bile duct paucity	93403	Synpolydactyly
3237	Symphalangism-brachydactyly syndrome	261619	Syndromic bile duct paucity due to a JAG1 point mutation	295195	Synpolydactyly type 1
465508	Symptomatic form of classic hemochromatosis	261629	Syndromic bile duct paucity due to a NOTCH2 point mutation	295197	Synpolydactyly type 2
276630	Symptomatic form of Coffin-Lowry syndrome in female carriers	261600	Syndromic bile duct paucity due to monosomy 20p12	295199	Synpolydactyly type 3
449291	Symptomatic form of fragile X syndrome in female carrier	84064	Syndromic diarrhea	295197	Synpolydactyly, Debeer type
465508	Symptomatic form of hemochromatosis type 1	84064	Syndromic diarrhea/Tricho-hepato-enteric syndrome	295199	Synpolydactyly, Malik type
177926	Symptomatic form of hemophilia A in female carriers	77298	Syndromic microphthalmia type 3	295195	Synpolydactyly, Vordingborg type
177929	Symptomatic form of hemophilia B in female carriers	85275	Syndromic microphthalmia type 4	3275	Synspondylism
465508	Symptomatic form of HFE-related hereditary hemochromatosis	178364	Syndromic microphthalmia type 5	93926	Syntelencephaly
206546	Symptomatic form of muscular dystrophy of Duchenne and Becker in female carriers	139471	Syndromic microphthalmia type 6	840	Syringadenoma papilliferum
357332	Synactyly-camptodactyly and clinodactyly of fifth fingers-bifid halluces syndrome	2556	Syndromic microphthalmia type 7	840	Syringocystadenoma papilliferum
98915	Synaptic congenital myasthenic syndromes	3434	Syndromic microphthalmia type 8	314701	Systemic AL amyloidosis
93404	Syndactyly of fingers 4 and 5	2470	Syndromic microphthalmia type 9	188	Systemic capillary leak syndrome
93402	Syndactyly type 1	77299	Syndromic microphthalmia type 10	→528	Systemic cystic angiomas-Seip syndrome
295193	Syndactyly type 1, Castilla type	178364	Syndromic microphthalmia/anophthalmia due to OTX2 mutation	364033	Systemic EBV+ T-cell LPD of childhood
295189	Syndactyly type 1, Lueken type	228426	Syndromic multisystem autoimmune disease due to Itch deficiency	364033	Systemic EBV-positive T-cell lymphoproliferative disease of childhood
295191	Syndactyly type 1, Montagu type	98606	Syndromic orbital border hypoplasia	364033	Systemic Epstein-Barr virus-positive T-cell lymphoproliferative disease of childhood
295187	Syndactyly type 1, Weidenreich type	281090	Syndromic recessive X-linked ichthyosis	401996	Systemic karyomegaly
295187	Syndactyly type 1a	281090	Syndromic RXLI	98849	Systemic mastocytosis with an associated clonal hematologic non-mast cell lineage disease
295189	Syndactyly type 1b	457223	Syndromic sensorineural deafness due to combined oxidative phosphorylation defect	90069	Systemic monochloroacetate poisoning
295191	Syndactyly type 1c	457223	Syndromic sensorineural deafness due to COXPD	439762	Systemic PAN
295193	Syndactyly type 1d	457223	Syndromic sensorineural hearing loss due to COXPD	439762	Systemic periarteritis nodosa
93403	Syndactyly type 2	281090	Syndromic X-linked ichthyosis	439762	Systemic polyarteritis nodosa
93404	Syndactyly type 3			85414	Systemic polyarthritis
93405	Syndactyly type 4			158	Systemic primary carnitine deficiency
93406	Syndactyly type 5			90291	Systemic scleroderma
295012	Syndactyly type 6			90291	Systemic sclerosis

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
220407	Systemic sclerosis sine scleroderma	66529	Takotsubo syndrome	2432	Teebi-Al Saleh-Hassoon syndrome
85414	Systemic-onset JIA	101028	TALDO	1094	Teebi-Kaurah syndrome
85414	Systemic-onset juvenile idiopathic arthritis	2886	Talipes equinovarus-atrial septal defect-Robin sequence-persistence of the left superior vena cava syndrome	1974	Teebi-Naguib-Alawadi syndrome
134	T2 deficiency			3291	Teebi-Shaltout syndrome
99861	T-ALL			3292	Tel Hashomer camptodactyly syndrome
169160	T-B+ SCID due to CD3delta/CD3epsilon/CD3zeta	217335	Tall forehead-sparse hair-skin hyperextensibility-scoliosis syndrome	90389	Telangiectasia macularis eruptiva perstans
169157	T-B+ SCID due to CD45 deficiency	404443	Tall stature-intellectual disability-facial dysmorphism syndrome	284227	Telangiectasia-erythrocytosis-monoclonal gammopathy-perinephric-fluid collections-intrapulmonary shunting syndrome
276	T-B+ SCID due to gamma chain deficiency				
169154	T-B+ SCID due to IL-7Ralpha deficiency	329191	Tall stature-scoliosis-macroductyly of the great toes syndrome	3293	Telecanthus-hypertelorism-strabismus-pes cavus syndrome
35078	T-B+ SCID due to JAK3 deficiency	329191	Tall stature-scoliosis-macroductyly of the halluces syndrome		
169160	T-B+ severe combined immunodeficiency due to CD3delta/CD3epsilon/CD3zeta	50809	Talo-patello-scapoid osteolysis	2885	Telfer-Sugar-Jaeger syndrome
169157	T-B+ severe combined immunodeficiency due to CD45 deficiency	31150	Tangier disease	1596	Telomeric 15q deletion syndrome
		180	Tapetochoroidal dystrophy	36367	Telomeric deletion 1q
		98839	Tappeiner-Pfleger disease	280	Telomeric deletion 4p
276	T-B+ severe combined immunodeficiency due to gamma chain deficiency	3320	TAR syndrome	96145	Telomeric deletion 4q
		65250	Tarlov cyst	1627	Telomeric deletion 5q
		2886	TARP syndrome	96126	Telomeric deletion 7p
169154	T-B+ severe combined immunodeficiency due to IL-7Ralpha deficiency	99170	Tarsal kink syndrome	1636	Telomeric deletion 7q36
		1412	Tarsal-carpal coalition syndrome	1642	Telomeric deletion 9p
		371	Tarui disease	1580	Telomeric deletion 10p
35078	T-B+ severe combined immunodeficiency due to JAK3 deficiency	163654	Tattoo dysplasia	96148	Telomeric deletion 10q
		2731	Taurodontia-absent teeth-sparse hair syndrome	2308	Telomeric deletion 11q
276	T-B+ severe combined immunodeficiency, X-linked	3289	Taurodontism	96149	Telomeric deletion 12q
		99045	Taussig-Bing syndrome	96150	Telomeric deletion 14q
86871	T-cell chronic lymphocytic leukemia	→33364	Tay syndrome	531	Telomeric deletion 17p
324294	T-cell immunodeficiency due to RHOH deficiency	845	Tay-Sachs disease	1597	Telomeric deletion 17q
		309239	Tay-Sachs disease, B1 variant	96129	Telomeric deletion 19p
324294	T-cell immunodeficiency with epidermodysplasia verruciformis	309192	Tay-Sachs disease, B variant, adult form	96152	Telomeric deletion 20q
		309178	Tay-Sachs disease, B variant, infantile form	1590	Telomeric deletion13q
86872	T-cell large granular lymphocyte leukemia	309185	Tay-Sachs disease, B variant, juvenile form	96069	Telomeric duplication 1p36
86872	T-cell LGL leukemia	669	Taybi syndrome	96070	Telomeric duplication 2p
86886	T-cell lymphoma, AILD type	90650	Taybi syndrome	96094	Telomeric duplication 2q
86871	T-cell prolymphocytic leukemia	2636	Taybi-Linder syndrome	96071	Telomeric duplication 3p
300857	T-cell/histiocyte rich large B cell lymphoma	98960	TBCD	96072	Telomeric duplication 4p
86872	T-LGL	857	TBS	96096	Telomeric duplication 4q
86871	T-PLL	2967	TCI deficiency	96097	Telomeric duplication 5q
1350	Tabatznik syndrome	103918	TCP	1745	Telomeric duplication 6p
3384	TAC	397959	TCR-alpha-beta+ T-cell deficiency	96098	Telomeric duplication 6q
447896	TACH syndrome	397959	TCR-alpha-beta-positive T-cell deficiency	96074	Telomeric duplication 7p
457077	TAFRO syndrome	2655	TD	96100	Telomeric duplication 8q
567	Takao syndrome	1860	TD1	96101	Telomeric duplication 9q
2905	Takatsuki syndrome	93274	TD2	96102	Telomeric duplication 10q
3287	Takayasu arteritis	3352	TDO syndrome	96103	Telomeric duplication 11q
66529	Tako-Tsubo cardiomyopathy	1519	Teebi hypertelorism syndrome	96105	Telomeric duplication 13q
66529	Tako-Tsubo syndrome	1519	Teebi syndrome	1705	Telomeric duplication 14q
66529	Takotsubo cardiomyopathy	1519	Teebi syndrome	1707	Telomeric duplication 15q
				96078	Telomeric duplication 16p
				96106	Telomeric duplication 16q
				3379	Telomeric duplication 17q
				1716	Telomeric duplication 18q

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
1717	Telomeric duplication 19q	3309	Tetrasomy 5p	49827	Thiamine-responsive megaloblastic anemia with diabetes mellitus and sensorineural deafness
96107	Telomeric duplication 20q	3310	Tetrasomy 9p		
96109	Telomeric duplication 22q	289522	Tetrasomy 11q24.1	268184	Thiamine-responsive MSUD
1762	Telomeric duplication Xq	884	Tetrasomy 12p		
1620	Telomeric monosomy 3p	314588	Tetrasomy 15(q25-qter)	2405	Thickened earlobes-conductive deafness syndrome
75565	TEMF	314588	Tetrasomy 15q26		
352737	Temperature-sensitive oculocutaneous albinism type 1	485405	Tetrasomy 16p12.1-p12.3	98960	Thiel-Behnke corneal dystrophy
		485405	Tetrasomy 16p12.1p12.3	3314	Thiemann disease, familial form
284227	TEMP1 syndrome	3307	Tetrasomy 18p	3235	Thies-Reis syndrome
420561	Temple-Baraitser syndrome	96055	Tetrasomy 21	1506	Thin ribs-tubular bones-dysmorphism syndrome
397	Temporal arteritis	9	Tetrasomy X		
363417	Temtamy preaxial brachydactyly syndrome	140917	Teunissen-Cremers syndrome	166424	Thinking seizures
		746	TFP deficiency	→300	Thiolase deficiency
1777	Temtamy syndrome	746	TFPD	141030	Third branchial cleft anomaly
1777	Temtamy-Shalash syndrome	225123	TFR2-related hemochromatosis	141030	Third branchial cleft cyst
66627	Tenosynovial giant cell tumor	476113	TFRC-related combined immunodeficiency	141030	Third branchial cleft fistula
137834	Ter Haar syndrome		TGA with cardiac malformation	363444	THOC6-related developmental delay-microcephaly-facial dysmorphism syndrome
252018	Teratoma of the central nervous system	216729	TGA with coarctation		
141107	Teratoma of the nasopharynx	99042	TGCT	2547	Thomas-Jewett-Raines syndrome
363483	Teratoma of the testis	66627	TH-SHFM	2031	Thompson-Baraitser syndrome
88630	Terminal osseous dysplasia-pigmentary defects syndrome	3329	TH-DONNAI syndrome	614	Thomsen and Becker disease
		1780	Thakker-Donnai syndrome	2866	Thong-Douglas-Ferrante syndrome
93937	Terminal transverse defects of arm	3312	Thalidomide embryopathy	1861	Thoracic dysplasia-hydrocephalus syndrome
141242	Tessier number 1 cleft	2655	Thanatophoric dwarfism		
141258	Tessier number 4 facial cleft	1860	Thanatophoric dwarfism type 1	97330	Thoracic outlet compression syndrome
141261	Tessier number 5 facial cleft	93274	Thanatophoric dwarfism type 2		
141265	Tessier number 6 facial cleft	93274	Thanatophoric dwarfism-cloverleaf skull syndrome	97330	Thoracic outlet syndrome
325124	Testicular agenesis	2655	Thanatophoric dysplasia	1759	Thoraco-abdominal enteric duplication
363494	Testicular non seminomatous germ cell tumor	1860	Thanatophoric dysplasia type 1		
363494	Testicular non-dysgerminomatous germ cell tumor	93274	Thanatophoric dysplasia type 2	1335	Thoraco-abdominal syndrome
983	Testicular regression syndrome	→175	Thanatophoric dysplasia, Glasgow variant	3317	Thoracolumbar pelvic dysplasia
842	Testicular seminoma	436169	THBD-related bleeding disorder	268384	Thoracolumbosacral spina bifida aperta
842	Testicular seminomatous germ cell tumor	436169	THBD-related coagulopathy	268752	Thoracolumbosacral spina bifida cystica
363489	Testicular sex cord-stromal tumor	99917	Theca (steroid-producing) cell cancer, not further specified		
363483	Testicular teratoma	99917	Theca steroid-producing cell malignant tumor of ovary, not further specified	1803	Thoracomelic dysplasia
3000	Testotoxicosis		99917	Theodore's superior limbic keratoconjunctivitis	→2199
3299	Tetanus	88633	Theodore's syndrome	300857	THRLBCL
9	Tetra X	88633	Thiamine-responsive BCKD deficiency	36258	Thromboangiitis obliterans
294971	Tetra-amelia		268184	Thiamine-responsive branched-chain 2-ketoacid dehydrogenase deficiency	3204
3301	Tetraamelia-multiple malformations syndrome	268184	Thiamine-responsive encephalopathy	67044	Thrombocytopenia with congenital dyserythropoietic anemia
199310	Tetragametic chimerism		199348	Thiamine-responsive maple syrup urine disease	3320
293284	Tetrahydrobiopterin-responsive HPA/PKU	268184	Thiamine-responsive megaloblastic anemia syndrome	457077	Thrombocytopenia-anasarca-renal insufficiency-organomegaly syndrome
293284	Tetrahydrobiopterin-responsive hyperphenylalaninemia/phenylketonuria		49827	Thiamine-responsive megaloblastic anemia syndrome	3323
3303	Tetralogy of Fallot	49827	Thiamine-responsive megaloblastic anemia syndrome	3002	Thrombocytopenic purpura, autoimmune
2564	Tetramelic monodactyly				
3305	Tetraploidy				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
436169	Thrombomodulin-related bleeding disorder	295077	Tibial longitudinal meromelia, unilateral	268748	Total spina bifida cystica
				2796	Touraine-Solente-Gole syndrome
436169	Thrombomodulin-related coagulopathy	609	Tibial muscular dystrophy	857	Townes syndrome
				295028	Tibio-fibular fusion
54057	Thrombotic thrombocytopenic purpura	295028	Tibio-fibular synostosis	95455	Toxic epidermal necrolysis
				294981	Tibiofibular terminal transverse meromelia
2251	Thumb deformity-alopecia-pigmentation anomaly syndrome	295099	Tibiofibular terminal transverse meromelia, bilateral	95455	Toxic epidermolysis
				279894	Toxic maculopathy due to antimalarial drugs
294988	Thumb hypodactyly	295097	Tibiofibular terminal transverse meromelia, unilateral	227972	Toxic oil syndrome
295112	Thumb hypodactyly, bilateral			293173	Toxic pustuloderma
295110	Thumb hypodactyly, unilateral	297	Tick-borne encephalitis	230800	Toxin-mediated infectious botulism
294988	Thumb oligodactyly			230800	Toxin-mediated infective botulism
295112	Thumb oligodactyly, bilateral	42665	Tietz syndrome	3343	Toxocariasis
295110	Thumb oligodactyly, unilateral	1662	Tight skin contracture syndrome	858	Toxoplasma embryofetopathy
1078	Thumb stiffness-brachydactyly-intellectual disability syndrome	65283	Timothy syndrome	858	Toxoplasma embryopathy
		91500	TINU syndrome	93164	TPHA
2919	Thurston syndrome	352540	TIO	444463	TPPII deficiency
83471	Thymic aplasia	480483	TJP2 deficit	444463	TPPII-related immunodeficiency, autoimmunity, and neurodevelopmental delay with impaired glycolysis and lysosomal expansion disease
99868	Thymic carcinoma	420561	TMBTS		
99869	Thymic neuroendocrine carcinoma	→1394	TMCO1 defect syndrome		
97289	Thymic neuroendocrine tumor	420611	TMD		
3326	Thymic-renal-anal-lung dysplasia	609	TMD		
99867	Thymoma	314667	TMEM165-CDG		
263310	Thymoma type A	466703	TMEM199-CDG	2950	TPT-PS syndrome
263324	Thymoma type AB	1194	TMEM70-related mitochondrial encephalo-cardio-myopathy	412022	Traboulsi syndrome
263317	Thymoma type B			3346	Tracheal agenesis
169105	Thymoma-immunodeficiency syndrome	99886	TNDM	2042	Tracheo-esophageal fistula-hypospadias syndrome
3327	Thyrocerbrorenal syndrome	32960	TNF receptor 1-associated periodic syndrome	3347	Tracheobronchomegaly
95716	Thyroid dysmorphogenesis	64686	Tolosa-Hunt syndrome	3348	Tracheobronchopathia osteochondroplastica
95712	Thyroid ectopia	1920	Toluene embryopathy	3348	Tracheopathia osteoplastica
95719	Thyroid hemigenesis	640	Tomaculous neuropathy	3052	Tranebjaerg-Svejgaard syndrome
95720	Thyroid hypoplasia	→314632	Tomé-Brunet-Fardeau syndrome	101028	Transaldolase deficiency
97285	Thyroid lymphoma	454718	Tonic pupil-tendon areflexia syndrome	859	Transcobalamin deficiency
91347	Thyroid stimulating hormone-secreting pituitary adenoma			2967	Transcobalamin I deficiency
2091	Thyroid-renal-digital anomalies	1547	Tonoki-Ohura-Niikawa syndrome	859	Transcobalamin II deficiency
79102	Thyrotoxic hypokalemic periodic paralysis	2228	Tooth and nail syndrome	2967	Transcobalamin-1 deficiency
		3460	Torg-Winchester syndrome	199247	Transcortin deficiency
79102	Thyrotoxic periodic paralysis	1827	Toriello syndrome	495	Transgrediens et progrediens palmoplantar keratoderma
91347	Thyrotroph adenoma	3338	Toriello-Carey syndrome	495	Transgrediens et progrediens PPK
2768	Tibia vara Blount	79347	Toriello-Higgins-Miller syndrome	87503	Transgrediens palmoplantar keratoderma of Siemens
3329	Tibial aplasia-ectrodactyly syndrome	3339	Toriello-Lacassie-Droste syndrome	420611	Transient abnormal myelopoiesis
93322	Tibial hemimelia	51084	Torsade-de-pointes syndrome with short coupling interval	98871	Transient acquired pure red cell aplasia
				3341	Torticollis-keloids-cryptorchidism-renal dysplasia syndrome
3329	Tibial hemimelia with split hand/foot malformation	75326	Tortuosity of retinal arteries	79411	Transient bullous dermolysis of the newborn
295079	Tibial hemimelia, bilateral	97330	TOS	98871	Transient erythroblastopenia of childhood
295077	Tibial hemimelia, unilateral	2701	Tosti syndrome		
3329	Tibial hemimelia-ectrodactyly syndrome	294971	Total amelia	2312	Transient familial neonatal hyperbilirubinemia
		49382	Total color blindness		
93322	Tibial longitudinal meromelia	98994	Total early-onset cataract	289877	Transient hyperammonemia of the newborn
295079	Tibial longitudinal meromelia, bilateral	180126	Total septate uterus		
		268377	Total spina bifida aperta		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
169139	Transient hypogammaglobulinemia of infancy	447896	Tremor-ataxia-central hypomyelination syndrome	→33364	Trichothiodystrophy-osteosclerosis syndrome
300293	Transient infantile hypertriglyceridemia and fatty liver	3350	Tremor-nystagmus-duodenal ulcer syndrome	→33364	Trichothiodystrophy-sun sensitivity syndrome
300293	Transient infantile hypertriglyceridemia and hepatosteatorrhea	64694	Trench fever	1209	Tricuspid atresia
		1822	Trevor disease	95457	Tricuspid valve agenesis
		99832	TRH resistance syndrome	95458	Tricuspid valve prolapse
66529	Transient left ventricular apical ballooning syndrome	2970	Triad syndrome	221091	Trigeminal neuralgia
		444463	TRIANGLE disease	98908	Triglyceride deposit cardiomyovasculopathy
420611	Transient myeloproliferative disease	85170	Triangular tibia-fibular aplasia syndrome		1308
420611	Transient myeloproliferative syndrome	863	Trichinellosis	3368	Trigonocephaly-bifid nose-acral anomalies syndrome
		863	Trichinosis		
391504	Transient neonatal acquired myasthenia	3352	Tricho-dento-osseous syndrome	3365	Trigonocephaly-broad thumbs syndrome
		84064	Tricho-hepato-enteric syndrome		
391504	Transient neonatal autoimmune myasthenia gravis	3354	Tricho-oculo-dermo-vertebral syndrome	3369	Trigonocephaly-short stature-developmental delay syndrome
280615	Transient neonatal cyanosis and anemia due to Toms River Hemoglobin	1264	Tricho-retino-dento-digital syndrome	401764	Trilineage bone marrow failure-developmental delay syndrome
		3351	Trichodental syndrome	3374	Triopia
99886	Transient neonatal diabetes mellitus	3360	Trichodermal syndrome-intellectual disability syndrome	868	Triose phosphate-isomerase deficiency
329942	Transient neonatal glutaric acidemia type 2	3353	Trichodermodyplasia-dental alterations syndrome	485405	Trip(16)(p12.1p12.3)
				444463	Tripeptidyl-peptidase II deficiency
329942	Transient neonatal glutaric aciduria type 2	228379	Trichodysplasia spinulosa	2950	Triphalangeal thumb-polysyndactyly syndrome
329942	Transient neonatal MAD deficiency	79129	Trichodysplasia-amelogenesis imperfecta syndrome	2947	Triphalangeal thumbs-brachyectrodactyly syndrome
329942	Transient neonatal MADD				
329942	Transient neonatal multiple acyl-CoA dehydrogenase deficiency	3361	Trichodysplasia-xeroderma syndrome	3133	Triphalangeal thumbs-dislocation of patella syndrome
391504	Transient neonatal myasthenia gravis	864	Trichofolliculoma	869	Triple A syndrome
		84064	Trichohepatoenteric syndrome		
93164	Transient pseudohypoaldosteronism	3363	Trichomegaly-retina pigmentary degeneration-dwarfism syndrome	415	Triple H syndrome
				3375	Triple X syndrome
3402	Transient tyrosinemia of the neonate	3355	Trichoodontoonychia dysplasia	3375	Triplo-X syndrome
3402	Transient tyrosinemia of the newborn	3355	Trichoodontoonychia dysplasia with bone deficiency in frontoparietal region	3376	Triploidy
213746	Transitional cell carcinoma of the corpus uteri	565	Trichopoliodystrophy	96069	Trisomy 1pter
		77258	Trichorhinopalangeal syndrome type 1 and 3	261344	Trisomy 1q
280224	Transitional PMD			250994	Trisomy 1q21.1
319308	Translocation renal cell carcinoma	502	Trichorhinopalangeal syndrome type 2	96070	Trisomy 2pter
85451	Transthyretin amyloid cardiopathy	→33364	Trichorrhexis nodosa syndrome	313947	Trisomy 2q23.1
85447	Transthyretin amyloid neuropathy	33364	Trichothiodystrophy	294026	Trisomy 2q31.1
85447	Transthyretin amyloid polyneuropathy	→33364	Trichothiodystrophy type B	96094	Trisomy 2qter
		→33364	Trichothiodystrophy type C	96071	Trisomy 3pter
85451	Transthyretin-related familial amyloid cardiomyopathy	→33364	Trichothiodystrophy type D	96095	Trisomy 3q26
		→33364	Trichothiodystrophy type E	251038	Trisomy 3q29
→221061	Transverse limb deficiency-hemangioma syndrome	→33364	Trichothiodystrophy type F	1738	Trisomy 4p
		→33364	Trichothiodystrophy type G	96072	Trisomy 4pter
180160	Transverse vaginal septum	→33364	Trichothiodystrophy type G	96096	Trisomy 4qter
32960	TRAPS syndrome	→33364	Trichothiodystrophy with congenital ichthyosis	1742	Trisomy 5p
399175	Traumatic avascular necrosis			329802	Trisomy 5p13
399175	Traumatic AVN			228415	Trisomy 5q35
861	Treacher-Collins syndrome	→33364	Trichothiodystrophy-neurocutaneous syndrome	96097	Trisomy 5qter
→1215	Treft-Sanborn-Carey syndrome			1745	Trisomy 6pter

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
96098	Trisomy 6qter	3375	Trisomy X	91500	Tubulointerstitial nephritis and uveitis syndrome
314034	Trisomy 7p22.1	217377	Trisomy Xp11.22-p11.23	2997	Tucker syndrome
96074	Trisomy 7pter	261483	Trisomy Xq27.3-q28	→2036	Tuffli-Laxova syndrome
96121	Trisomy 7q11.23	261483	Trisomy Xq27.3q28	1063	Tufted angioma
264450	Trisomy 8p	1762	Trisomy Xq28	3392	Tularemia
251076	Trisomy 8p23.1	88629	Tritan colour blindness	640	Tulip-bulb digger's palsy
1752	Trisomy 8q	88629	Tritanopia	32960	Tumor necrosis factor receptor 1 associated periodic syndrome
228399	Trisomy 8q12	49827	TRMA	289539	Tumor susceptibility linked to germline BAP1 mutations
96100	Trisomy 8qter	1349	tRNA-LYS-related cardiomyopathy-hearing loss syndrome	352540	Tumor-induced osteomalacia
236	Trisomy 9p	103918	Tropical calcific chronic pancreatitis	879	Tungiasis
96101	Trisomy 9qter	75565	Tropical endomyocardial fibrosis	3225	Tungland-Bellman syndrome
171929	Trisomy 10p	99654	Tropical pancreatic diabetes	99053	Tunnel subaortic stenosis
276422	Trisomy 10q22.3q23.3	103918	Tropical pancreatitis	211	Turban tumor syndrome
96102	Trisomy 10qter	764	Tropical pyomyositis	99818	Turcot syndrome with polyposis
300305	Trisomy 11p15.4	289326	Tropical spastic paraparesis	881	Turner syndrome
96103	Trisomy 11qter	101000	Troyer syndrome	99413	Turner syndrome due to structural X chromosome anomalies
1699	Trisomy 12p	983	TRS	2614	Turner-Kieser syndrome
3378	Trisomy 13	313906	True congenital pancreatic cyst	63440	Turriccephaly
96105	Trisomy 13qter	2138	True hermaphroditism	79153	Twenty-nail dystrophy
261229	Trisomy 14q11.2	2512	True microcephaly	95431	Twin to twin transfusion syndrome
1705	Trisomy 14qter	180074	True unicornuate uterus	1461	Twisted atrioventricular connections
238446	Trisomy 15q11-q13	3357	Trueb-Burg-Bottani syndrome	2889	Twisted hair
238446	Trisomy 15q11q13	3384	Truncus arteriosus	2198	Tylosis-oesophageal carcinoma syndrome
1707	Trisomy 15qter	228379	TS	477781	Type 1 condylar hyperplasia
261204	Trisomy 16p11.2p12.2	352737	TS OCA type 1	3255	Type 1 syndactyly-microcephaly-intellectual disability syndrome
261243	Trisomy 16p13.11	3173	Tsao-Ellingson syndrome	→79259	Type 1C glycogenosis
96078	Trisomy 16pter	66627	TSGCT	→79259	Type 1D glycogenosis
96106	Trisomy 16qter	91347	TSH-oma	93554	Type II mixed cryoglobulinemia
261290	Trisomy 17p	91347	TSH-secreting pituitary adenoma	99745	Typhoid
1713	Trisomy 17p11.2	289326	TSP	99745	Typhoid fever
477817	Trisomy 17p11.2-p12	3268	Tsukahara syndrome	99745	Typhoidal salmonellosis
477817	Trisomy 17p11.2p12	3387	Tsukahara-Kajii syndrome	90038	Typical hemolytic-uremic syndrome
217385	Trisomy 17p13.3	83317	Tsutsugamushi disease	90038	Typical HUS
139474	Trisomy 17q11.2	83317	Tsutsugamushi fever	171436	Typical nemaline myopathy
261272	Trisomy 17q12	54057	TTP	158766	Typical urticaria pigmentosa
217340	Trisomy 17q21.31	85447	TTR amyloid neuropathy	1895	Typus Edinburgensis
3379	Trisomy 17qter	85451	TTR-related amyloid cardiomyopathy	79431	Tyrosinase-negative oculocutaneous albinism
3380	Trisomy 18	85451	TTR-related amyloid cardiac amyloidosis	101150	Tyrosine hydroxylase deficiency
1715	Trisomy 18p	180242	Tubal cancer	101150	Tyrosine hydroxylase-deficient dopa-responsive dystonia
1716	Trisomy 18qter	3389	Tuberculosis	69723	Tyrosinemia due to 4-hydroxyphenylpyruvate dioxygenase deficiency
1717	Trisomy 19qter	805	Tuberous sclerosis	69723	Tyrosinemia due to 4-hydroxyphenylpyruvic acid oxidase deficiency
261318	Trisomy 20p	805	Tuberous sclerosis complex	69723	Tyrosinemia due to HPD deficiency
96107	Trisomy 20qter	88924	Tuberous sclerosis/polycystic kidney disease contiguous gene syndrome		
870	Trisomy 21	2593	Tubular aggregate myopathy		
1727	Trisomy 22q11.2	100048	Tubular duplication of the esophagus		
96109	Trisomy 22qter	73224	Tubular renal disease-cardiomyopathy syndrome		
1738	Trisomy of the short arm of chromosome 4	467166	Tubulinopathy-associated dysgyria		
1742	Trisomy of the short arm of chromosome 5	319325	Tubulocystic renal cell carcinoma		
236	Trisomy of the short arm of chromosome 9				
1715	Trisomy of the short arm of chromosome 18				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
28378	Tyrosinemia due to TAT deficiency	98825	Unclassified mixed myelodysplastic/myeloproliferatic syndrome	97363	Unilateral multicystic dysplastic kidney
28378	Tyrosinemia due to tyrosine aminotransferase deficiency			97363	Unilateral multicystic renal dysplasia
882	Tyrosinemia type 1	98827	Unclassified myelodysplastic syndrome	268943	Unilateral polymicrogyria
28378	Tyrosinemia type 2			295148	Unilateral PPD2
69723	Tyrosinemia type 3	98825	Unclassified myelodysplastic/myeloproliferative disease	295012	Unilateral syndactyly of digits 2-5
882	Tyrosinemia type I			1464	Univentricular heart
28378	Tyrosinemia type II	251328	Unclassified vasculitis	99069	Univentricular heart with single atrio-ventricular valve
69723	Tyrosinemia type III				
75840	UCMD	1410	Uncombable hair syndrome	79146	Universal melanosis
90002	UCTD	1264	Uncombable hair-retinal pigmentary dystrophy-dental anomalies-brachydactyly syndrome	620	Universal mesentery
609	Udd myopathy			84096	Unknown leukodystrophy
79238	UDP-galactose-4-epimerase deficiency	103920	Undetermined colitis	99104	Unroofed coronary sinus
178315	UES	442835	Undetermined early-onset epileptic encephalopathy	91140	Unspecified JIA
205	UGT deficiency	442835	Undetermined EOEE	91140	Unspecified juvenile idiopathic arthritis
79234	UGT deficiency type 1	418951	Undifferentiated carcinoma of esophagus	99139	Unstable hemoglobin disease
79235	UGT deficiency type 2			308	Unverricht-Lundborg disease
3403	Uhl anomaly	424970	Undifferentiated carcinoma of liver and IBT	251009	UPD(1)mat
2032	UIP			251004	UPD(1)pat
3404	Ulbright-Hodes syndrome	424970	Undifferentiated carcinoma of liver and intrahepatic biliary tract	96179	UPD(2)mat
308	ULD			96180	UPD(4)mat
3406	Ulerythema ophryogenesis	424080	Undifferentiated carcinoma of pancreas with osteoclast-like giant cells	96190	UPD(5)pat
320	Ulick syndrome			96181	UPD(6)mat
75840	Ullrich disease	423786	Undifferentiated carcinoma of stomach	96191	UPD(6)pat
2497	Ulna hypoplasia			96182	UPD(7)mat
2249	Ulna hypoplasia-intellectual disability syndrome	213721	Undifferentiated carcinoma of the corpus uteri	96192	UPD(7)pat
1837	Ulna metaphyseal dysplasia syndrome			96183	UPD(9)mat
93320	Ulnar clubhand	90002	Undifferentiated connective tissue syndrome	231147	UPD(11)mat
93320	Ulnar hemimelia	178315	Undifferentiated embryonal sarcoma of the liver	96193	UPD(11)pat
295073	Ulnar hemimelia, bilateral			97678	UPD(13)mat
295075	Ulnar hemimelia, unilateral	418951	Undifferentiated esophageal carcinoma	99324	UPD(13)pat
1122	Ulnar hypoplasia-lobster-claw deformity of feet syndrome			96184	UPD(14)mat
		423786	Undifferentiated gastric carcinoma	96334	UPD(14)pat
1122	Ulnar hypoplasia-split foot syndrome	86830	Undifferentiated myeloproliferative disease	98754	UPD(15)mat
				2023	Undifferentiated pleomorphic sarcoma
93320	Ulnar longitudinal meromelia	178315	Undifferentiated sarcoma of the liver	96185	UPD(16)mat
295073	Ulnar longitudinal meromelia, bilateral			96186	UPD(20)mat
295075	Ulnar longitudinal meromelia, unilateral	251332	Unexplained long-lasting fever/inflammatory syndrome	96194	UPD(20)pat
				83468	Unicameral bone cyst
3138	Ulnar-mammary syndrome	180079	Unicornuate uterus with rudimentary horn	96195	UPD(21)pat
52056	Ulnar/fibula ray defect-brachydactyly syndrome			261519	UPD(X)mat
3405	Umbilical cord ulceration-intestinal atresia syndrome	180074	Unicornuate uterus without rudimentary horn	261524	UPD(X)pat
				3408	Upington disease
209886	UMOD-associated familial juvenile hyperuricemic nephropathy	93176	Unilateral congenital megacalycosis	2489	Upper limb defect-eye and ear abnormalities syndrome
209886	UMOD-associated FJHN	268947	Unilateral focal polymicrogyria	295049	Upper limb hypertrophy
35120	UMPH1 deficiency	101071	Unilateral hemispheric polymicrogyria	2497	Upper limb mesomelic dysplasia
3138	UMS			268740	Upper thoracic spina bifida aperta
104078	Unclassified intestinal pseudoobstruction	97363	Unilateral MCDK	268770	Upper thoracic spina bifida cystica
		99802	Unilateral megalencephaly	2023	UPS
				481665	UPS18 deficiency

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93583	Upshaw-Schulman syndrome	65681	Vaginal atresia	1053	Vein of Galen arteriovenous malformations
488	Urachal cyst	180247	Vaginal carcinoma		
431347	Urachal diverticulum	206489	Vaginal germ cell cancer	3424	Velo-facial-skeletal syndrome
431344	Urachal sinus	206489	Vaginal germ cell malignant tumor	567	Velocardiofacial syndrome
530	Urbach-Wiethe disease	180247	Vaginal malignant epithelial tumor	29207	Venereal arthritis
221145	Urban-Rifkin-Davis syndrome	158048	VAHS	319234	Venezuelan hemorrhagic fever
3409	Urban-Rogers-Meyer syndrome	88639	Valine metabolic defect	357131	Venous cervical rib syndrome
1839	Urban-Schosser-Spohn syndrome	228123	Valley fever	357131	Venous costoclavicular syndrome
94059	Uremic pruritus	99054	Valvular pulmonary stenosis	357131	Venous hyperabduction syndrome
105	Urethral atresia	1548	Van Benthem-Driessen-Hanveld syndrome	83454	Venous malformations with glomus cells
35120	Uridine 5'-monophosphate hydrolase deficiency		2806		
79238	Uridine diphosphate galactose-4-epimerase deficiency	2806	Van Bogaert encephalitis	357131	Venous thoracic outlet compression syndrome
30	Uridine monophosphate synthetase deficiency	3416	Van Buchem disease		
		1122	Van den Berghe-Dequecker syndrome	357131	Venous thoracic outlet syndrome
210128	Urocanic aciduria	3417	Van den Bosch syndrome	3201	Ventricular extrasystoles with syncopal episodes-perodactyly-Robin sequence syndrome
2704	Urofacial syndrome	2460	Van den Ende-Gupta syndrome		
83628	Urorectal septum malformation sequence	216796	Van der Hoeve syndrome	216694	Ventricular inversion
98606	Urrrets-Zavalia syndrome	2478	Van der Knaap syndrome		
79457	Urticaria pigmentosa	888	Van der Woude syndrome	99094	Ventricular septal defect with aortic insufficiency
886	USH	3419	Van Maldergem syndrome	216694	Ventriculoarterial and atrioventricular discordance
231169	USH1		Van Regemorter-Pierquin-Vamos syndrome		
231178	USH2	73	Vanishing bone disease	860	Ventriculoarterial discordance with atrioventricular concordance
231183	USH3	983	Vanishing testes syndrome		
886	Usher syndrome	983	Vanishing testis syndrome	443988	Ventriculomegaly-cystic kidney disease
231169	Usher syndrome type 1	729	Vaquez disease		
231178	Usher syndrome type 2	2754	Váradi syndrome	2899	Verloes-Bourguignon syndrome
231183	Usher syndrome type 3	2754	Váradi-Papp syndrome	2496	Verloes-David syndrome
2032	Usual interstitial pneumonia	454742	Variably protease-sensitive prionopathy	50817	Verloes-Deprez syndrome
213610	Uterine carcinosarcoma		Variant ABeta2M amyloidosis	2983	Verloes-Gillerot-Fryns syndrome
180145	Uterine cervical aplasia and agenesis	314652	Variant phenylketonuria	2551	Verloes-Van Maldergem-de Marneffe syndrome
180139	Uterine hypoplasia	79253	Variant PKU	3429	Verloove Vanhorick-Brubakk syndrome
439167	Uteroplacental vascular insufficiency	79473	Variagate porphyria	70476	Vernal keratoconjunctivitis
		404553	Vasculitis due to ADA2 deficiency		
180118	Uterus arcuatus	353356	Vasoproliferative tumor of ocular fundus	97282	Verner-Morrison syndrome
180118	Uterus cordiformis		Vasoproliferative tumor of retina	464318	Verrucous hemangioma
180129	Uterus subseptus	353356	Vasquez-Hurst-Sotos syndrome	79467	Verrucous nevus
178338	UV-sensitive syndrome	→261483	Västerbotten dystrophy	26793	Very long chain acyl-CoA dehydrogenase deficiency
1473	Uveal coloboma-cleft lip and palate-intellectual disability	85128	VATER association	431347	Vesicourachal diverticulum
39044	Uveal melanoma	887	Vater-like syndrome with pulmonary hypertension, abnormal ears and growth deficiency	252175	Vestibular schwannoma
3437	Uveomenigitic syndrome	52047	VATS	892	VHL
99771	Uvular cleft		VACTERL association	1493	Vici syndrome
370109	v-AT	228379	VCAN-related vitreoretinopathy	3433	Viljoen-Kallis-Voges syndrome
887	VACTERL association	898	VDDI	3434	Viljoen-Smart syndrome
3412	VACTERL with hydrocephalus	289157	VDDR II	97282	VIP-secreting tumor
887	VACTERL/VATER association	93160	VDDR-I	97282	VIPoma
25980	Vacuolar myopathy	289157	VDEGS	206991	Viral myositis
2478	Vacuolating megalencephalic leukoencephalopathy with subcortical cysts	2460	VDRR II	180176	Virginal breast hypertrophy
		93160	Vein of Galen aneurysm	99916	Virilizing ovarian tumor
		1053		158048	Virus-associated hemophagocytis syndrome

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228379	Virus-associated trichodysplasia spinulosa	2451	VMCM	897	Waardenburg syndrome type 4
		83454	VMGLOM	894	Waardenburg syndrome type I
280068	Visceral calciphylaxis	79124	VODI syndrome	895	Waardenburg syndrome type II
1876	Visceral myopathy-familial external ophthalmoplegia syndrome	3437	Vogt-Koyanagi-Harada disease	896	Waardenburg syndrome type III
		494	Vohwinkel syndrome	896	Waardenburg syndrome with limb anomalies
73246	Visceral neuropathy-brain anomalies-facial dysmorphism-developmental delay syndrome	79395	Vohwinkel syndrome with ichthyosis	897	Waardenburg-Hirschsprung syndrome
		2427	Volcke-Soekarman syndrome		
353344	Visible and exudative idiopathic juxtafoveal retinal telangiectasis	35737	Volubilis syndrome	98960	Waardenburg-Jonker corneal dystrophy
		83600	Von Economo encephalitis		
420556	Visual snow syndrome	364	Von Gierke disease	897	Waardenburg-Shah syndrome
3006	Vitamin B6-dependent seizures	98941	Von Hippel anomaly	280558	WABS
28	Vitamin B12-responsive methylmalonic acidemia	892	Von Hippel-Lindau disease	466943	WAC-related facial dysmorphism-developmental delay-behavioral abnormalities syndrome
		892	Von Hippel-Lindau syndrome		
79310	Vitamin B12-responsive methylmalonic acidemia type cblA	238557	Von Hippel-Lindau-dependent polycythemia	247709	Wagenmann-Froboese syndrome
79311	Vitamin B12-responsive methylmalonic acidemia type cblB	386	Von Meyenburg complexes disease	898	Wagner disease
		636	Von Recklinghausen disease	898	Wagner syndrome
308442	Vitamin B12-responsive methylmalonic acidemia, type cblDv2	363700	Von Recklinghausen disease due to NF1 mutation or intragenic deletion	893	WAGR syndrome
				90033	wAHA
28	Vitamin B12-responsive methylmalonic aciduria	3439	Von Voss-Cherstvoy syndrome	357332	Wahab syndrome
		903	Von Willebrand disease	90033	wAIHA
79310	Vitamin B12-responsive methylmalonic aciduria type cblA	166078	Von Willebrand disease type 1	2379	Waisman syndrome
		166081	Von Willebrand disease type 2	33226	Waldenström macroglobulinemia
79311	Vitamin B12-responsive methylmalonic aciduria, type cblB	166084	Von Willebrand disease type 2A	90362	Waldmann disease
		166087	Von Willebrand disease type 2B	1068	Walker-Dyson syndrome
308442	Vitamin B12-responsive methylmalonic aciduria, type cblDv2	166090	Von Willebrand disease type 2M	899	Walker-Warburg syndrome
		166093	Von Willebrand disease type 2N	1453	Wallis-Zieff-Goldblatt syndrome
27	Vitamin B12-unresponsive methylmalonic acidemia	166096	Von Willebrand disease type 3	2510	WARBM
		466934	VPS11-related autosomal recessive hypomyelinating leukodystrophy	2510	Warburg micro syndrome
79312	Vitamin B12-unresponsive methylmalonic acidemia type mut-	466934	VPS11-related autosomal recessive hypomyelinating leukoencephalopathy	3214	Warburg-Thomsen syndrome
				1052	Warburton-Anyane-Yeboah syndrome
289916	Vitamin B12-unresponsive methylmalonic acidemia type mut0	369852	VPS45 deficiency	96061	Warkany syndrome
27	Vitamin B12-unresponsive methylmalonic aciduria	353356	VPTR	90033	Warm AIHA
		99094	VSD with aortic insufficiency	1541	Warman-Mulliken-Hayward syndrome
79312	Vitamin B12-unresponsive methylmalonic aciduria type mut-	357131	VTOS	280558	Warsaw breakage syndrome
		137583	Vulvar intraepithelial neoplasia	51636	Warts-hypogammaglobulinemia-infections-myelokathexis syndrome
137583	Vulvar intraepithelial tumor				
289157	Vitamin D dependent rickets type I	83453	Vulvovaginal gingival syndrome	51636	Warts-infections-leukopenia-myelokathexis syndrome
289157	Vitamin D-dependency type I	206492	Vulvovaginal rhabdomyosarcoma	69745	Warty dyskeratoma
93160	Vitamin D-dependent rickets type II	53696	Vuopala disease	906	WAS
93160	Vitamin D-resistant rickets type II	888	VWS	1046	Water-West syndrome
1914	Vitamin K antagonists embryofetopathy	2804	W syndrome	100067	Waterhouse-Friderichsen syndrome
		2180	Waalder-Aarskog syndrome		
1243	Vitelliform macular dystrophy type 2	1106	Waardenburg anophthalmia syndrome	97282	Watery diarrhea-hypokalemia-achlorhydria syndrome
		3440	Waardenburg syndrome	→636	Watson syndrome
898	Vitreoretinal degeneration, Wagner type	894	Waardenburg syndrome type 1	33577	WCD
		895	Waardenburg syndrome type 2	284395	W DFA
26793	VLCAD deficiency	352740	Waardenburg syndrome type 2 with ocular albinism	97282	WDHA syndrome
26793	VLCADD			99971	W DLS
386	VMC	896	Waardenburg syndrome type 3	603	WDM
443988	VMCKD				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
3447	Weaver syndrome	171723	White sponge nevus	1409	Woolly hair-hypotrichosis-everted lower lip-outstanding ears syndrome
→3447	Weaver-like syndrome	171723	White sponge nevus of Cannon		
3448	Weaver-Williams syndrome	1489	Whooping cough	420686	Woolly hair-palmoplantar hyperkeratosis syndrome
33577	Weber-Christian disease	2779	Whyte-Murphy syndrome	65282	Woolly hair-palmoplantar hyperkeratosis-dilated cardiomyopathy syndrome
33577	Weber-Christian panniculitis	3454	Wieacker-Wolff syndrome		
1521	Webster-Deming syndrome	116	Wiedemann-Beckwith syndrome	420686	Woolly hair-palmoplantar keratoderma syndrome
900	Wegener granulomatosis	2156	Wiedemann-Oldigs-Oppermann syndrome		
228254	Weidman juvenile elastoma		3455	Wiedemann-Rautenstrauch syndrome	65282
3449	Weill-Marchesani syndrome				
3344	Weismann-Netter syndrome	319182	Wiedemann-Steiner syndrome	170	Woolly hair
3450	Weissenbacher- Zweymuller syndrome	85446	Wild type ABeta2-microglobulinic amyloidosis		
284395	Well-differentiated fetal adenocarcinoma of the lung	85446	Wild type ABeta2M amyloidosis	79414	Woolly hair nevus
99971	Well-differentiated liposarcoma	330001	Wild type ATTR amyloidosis	1409	Woolly hair-hypotrichosis-everted lower lip-outstanding ears syndrome
213736	Well-differentiated neuroendocrine neoplasm of the endometrium	330001	Wild type ATTR-related amyloidosis	65282	Woolly hair-palmoplantar hyperkeratosis-dilated cardiomyopathy syndrome
213736	Well-differentiated neuroendocrine tumor of the corpus uteri	3456	Wildervanck syndrome		
213736	Well-differentiated neuroendocrine tumor of the endometrium	739	Willi-Prader syndrome	65282	Woolly hair-palmoplantar keratoderma-dilated cardiomyopathy syndrome
		904	Williams syndrome		
263331	Well-differentiated thymic neuroendocrine carcinoma	904	Williams-Beuren syndrome	166277	Wormian bone-multiple fractures-dentinogenesis imperfecta-skeletal dysplasia
		411501	Williams-Campbell syndrome		
146	Well-differentiated thyroid carcinoma	51636	WILM	3465	Worster-Drought syndrome
1373	Wellesley-Carman-French syndrome	654	Wilms tumor	2790	Worth syndrome
901	Wells syndrome	220	Wilms tumor and pseudohermaphroditism	178475	Wound botulism
2815	Wells-Jankovic syndrome		893	Wilms tumor-aniridia-genitourinary anomalies-intellectual disability syndrome	165955
83330	Werdnig-Hoffmann disease	905	Wilson disease	2834	Wrinkled skin syndrome
652	Wermer syndrome	3459	Wilson-Turner syndrome	2834	Wrinkly skin syndrome
3332	Werner mesomelic syndrome	3460	Winchester syndrome	1667	WRS
902	Werner syndrome	169095	Winged helix deficiency	902	WS
1979	Werner-like syndrome due to combined growth factor deficiency	94087	Winkelmann cytophagic panniculitis	894	WS1
		2515	Winship-Viljoen-Leary syndrome	895	WS2
3451	West syndrome	906	Wiskott-Aldrich syndrome	896	WS3
83476	West-Nile encephalitis	829	Wissler-Fanconi syndrome	897	WS4
83476	West-Nile fever	2228	Witkop syndrome	163746	WS4 plus
2435	Westerhof-Beemer-Cormane syndrome	101068	Witschel dystrophy	2834	WSS
		→280	Wittwer syndrome	3466	WT limb-blood syndrome
83593	Western equine encephalitis	3237	WL syndrome	3459	WTS
83593	Western equine encephalomyelitis	247768	WNT4 deficiency	3411	Wunderlich syndrome
681	Westphall disease	1667	Wolcott-Rallison syndrome	899	WWS
952	Weyers acrofacial dysostosis	280	Wolf-Hirschhorn syndrome	53719	Wyburn-Mason syndrome
952	Weyers acrofacial dysostosis	3080	Wolff-Zimmermann syndrome	96201	X small rings
→2750	Whelan syndrome	3463	Wolfram syndrome	43	X-ALD
51636	WHIM syndrome	411590	Wolfram-like syndrome	300373	X-LAG (X-linked acrogigantism)
3452	Whipple disease	75233	Wolman disease	448348	X-LAG (X-linked acrogigantism) due to a point mutation
2053	Whistling face syndrome	3464	Woodhouse-Sakati syndrome	448372	X-LAG (X-linked acrogigantism) due to dup(X)q(26)
228290	White fibrous papulosis of the neck	2571	Woods-Black-Norbury syndrome	2182	X-linked aqueductal stenosis
2475	White forelock with malformations	137658	Woods-Crouchman-Huson syndrome		
3207	White matter hypoplasia-corpora callosa agenesis-intellectual disability syndrome		170	Woolly hair	448348
		79414	Woolly hair nevus		
370131	White platelet syndrome				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
448372	X-linked acrogigantism due to Xq26 microduplication	431140	X-linked colobomatous microphthalmia-microcephaly-short stature-psychomotor retardation syndrome	98863	X-linked Emery-Dreifuss muscular dystrophy
43	X-linked adrenoleukodystrophy			293621	X-linked endothelial corneal dystrophy
47	X-linked agammaglobulinemia			85294	X-linked epilepsy-learning disabilities-behavior disorders syndrome
43	X-linked ALD				
88917	X-linked Alport syndrome	1497	X-linked complicated corpus callosum dysgenesis	443197	X-linked erythropoietic protoporphyria
85278	X-linked Angelman-like syndrome	306617	X-linked complicated spastic paraplegia type 1	480880	X-linked facial dysmorphism-short stature-choanal atresia-intellectual disability syndrome limited to females
181	X-linked anhidrotic ectodermal dysplasia	90001	X-linked cone dysfunction syndrome with myopia		
85297	X-linked ataxia-deafness syndrome	95702	X-linked congenital adrenal hypoplasia	480880	X-linked female restricted facial dysmorphism-short stature-choanal atresia-intellectual disability
85292	X-linked ataxia-dementia syndrome	67044	X-linked congenital dyserythropoietic anemia with thrombocytopenia		
139583	X-linked auditory neuropathy with peripheral sensory neuropathy type 1	79495	X-linked congenital generalized hypertrichosis	→994	X-linked fetal akinesia syndrome
1131	X-linked branchial arch syndrome	565	X-linked copper deficiency	139583	X-linked hereditary sensory and autonomic neuropathy with deafness
481	X-linked BSMA	1661	X-linked corneal dermoid		
481	X-linked bulbospinal amyotrophy	52503	X-linked creatine transporter deficiency	139583	X-linked HSAN with deafness
391327	X-linked calvarial hyperostosis	85453	X-linked cutaneous amyloidosis	2182	X-linked HSAS
111	X-linked cardioskeletal myopathy and neutropenia	198	X-linked cutis laxa	2182	X-linked hydrocephalus
329235	X-linked central congenital hypothyroidism with late-onset macroorchidism	383	X-linked deafness type 2	2182	X-linked hydrocephalus with stenosis of aqueduct of Sylvius
		85321	X-linked deafness-intellectual disability syndrome	1397	X-linked hydrocephalus-cerebellar agenesis-intellectual disability syndrome
329235	X-linked central congenital hypothyroidism with late-onset testicular enlargement	139557	X-linked dHMN type 3	101088	X-linked hyper-IgM syndrome
596	X-linked centronuclear myopathy	1018	X-linked diffuse leiomyomatosis-Alport syndrome	181	X-linked hypohidrotic ectodermal dysplasia
139396	X-linked cerebral adrenoleukodystrophy	1145	X-linked distal arthrogyriposis multiplex congenita	89936	X-linked hypophosphatemia
163961	X-linked cerebral-cerebellar-coloboma syndrome	139557	X-linked distal hereditary motor neuropathy type 3	89936	X-linked hypophosphatemic rickets
101075	X-linked Charcot-Marie-Tooth disease type 1	139557	X-linked distal spinal muscular atrophy type 3	461	X-linked ichthyosis
101076	X-linked Charcot-Marie-Tooth disease type 2	35173	X-linked dominant chondrodysplasia punctata	231692	X-linked IGHD
101077	X-linked Charcot-Marie-Tooth disease type 3	163966	X-linked dominant chondrodysplasia, Chassaing-Lacombe type	317476	X-linked immunodeficiency with magnesium defect, Epstein-Barr virus infection and neoplasia
101078	X-linked Charcot-Marie-Tooth disease type 4	163966	X-linked dominant chondrodysplasia-hydrocephaly-microphthalmia syndrome	2571	X-linked immunoneurologic disorder
99014	X-linked Charcot-Marie-Tooth disease type 5			16	X-linked incomplete achromatopsia
352675	X-linked Charcot-Marie-Tooth disease type 6	443197	X-linked dominant erythropoietic protoporphyria	364028	X-linked intellectual disability due to GRIA3 anomalies
35173	X-linked chondrodysplasia punctata type 2	93951	X-linked dominant intellectual disability-epilepsy syndrome	3242	X-linked intellectual disability due to PQBP1 mutations
324601	X-linked cleft palate and ankyloglossia	443197	X-linked dominant protoporphyria	67045	X-linked intellectual disability with isolated growth hormone deficiency
		139557	X-linked dSMA type 3	776	X-linked intellectual disability with marfanoid habitus
431140	X-linked colobomatous microphthalmia-microcephaly-intellectual disability-short stature syndrome	363727	X-linked dyserythropoietic anemia with abnormal platelets and neutropenia	85273	X-linked intellectual disability, Abidi type
		373	X-linked dysplasia gigantism syndrome	85274	X-linked intellectual disability, Ahmad type
		53351	X-linked dystonia-parkinsonism	85276	X-linked intellectual disability, Armfield type
		75497	X-linked Ehlers-Danlos syndrome		

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
1193	X-linked intellectual disability, Atkin type	85326	X-linked intellectual disability, Stoll type	85282	X-linked intellectual disability-epileptic seizures-hypogenitalism-microcephaly-obesity syndrome
3056	X-linked intellectual disability, Brooks type	93950	X-linked intellectual disability, Sutherland-Haan type	480907	X-linked intellectual disability-global development delay-facial dysmorphism-sacral caudal remnant syndrome
85293	X-linked intellectual disability, Cabezas type	85328	X-linked intellectual disability, Turner type	3459	X-linked intellectual disability-gynecomastia-obesity syndrome
85277	X-linked intellectual disability, Cantagrel type	163976	X-linked intellectual disability, Van Esch type	85317	X-linked intellectual disability-hypogammaglobulinemia-progressive neurological deterioration syndrome
163971	X-linked intellectual disability, Cilliers type	→85293	X-linked intellectual disability, Vitale type	3055	X-linked intellectual disability-hypogonadism-ichthyosis-obesity-short stature syndrome
→93950	X-linked intellectual disability, Fichera type	85290	X-linked intellectual disability, Wilson type	59	X-linked intellectual disability-hypotonia syndrome
93947	X-linked intellectual disability, Golabi-Ito-Hall type	→280	X-linked intellectual disability, Wittwer type	85329	X-linked intellectual disability-hypotonia-facial dysmorphism-aggressive behavior syndrome
→457240	X-linked intellectual disability, Gu type	→59	X-linked intellectual disability, Zorick type	457260	X-linked intellectual disability-hypotonia-movement disorder syndrome
93952	X-linked intellectual disability, Hedera type	85327	X-linked intellectual disability-acromegaly-hyperactivity syndrome	→1762	X-linked intellectual disability-hypotonia-recurrent infections syndrome
163961	X-linked intellectual disability, Kroes type	85338	X-linked intellectual disability-ataxia-apraxia syndrome	423479	X-linked intellectual disability-limb spasticity-retinal dystrophy-diabetes insipidus syndrome
→1762	X-linked intellectual disability, Lubs type	324410	X-linked intellectual disability-cardiomegaly-congestive heart failure syndrome	85320	X-linked intellectual disability-macrocephaly-macroorchidism syndrome
85283	X-linked intellectual disability, Miles-Carpenter type	137831	X-linked intellectual disability-cerebellar hypoplasia syndrome	251383	X-linked intellectual disability-microcephaly-cortical malformation-thin habitus syndrome
163937	X-linked intellectual disability, Najm type	459070	X-linked intellectual disability-cerebellar hypoplasia-spondylo-epiphyseal dysplasia syndrome	163937	X-linked intellectual disability-microcephaly-pontocerebellar hypoplasia syndrome
163956	X-linked intellectual disability, Nascimento type	85295	X-linked intellectual disability-choreoathetosis-abnormal behavior syndrome	163971	X-linked intellectual disability-microcephaly-testicular failure syndrome
85322	X-linked intellectual disability, Pai type	85330	X-linked intellectual disability-corpora callosa agenesis-spastic quadriplegia syndrome	→3057	X-linked intellectual disability-monoamine oxidase A metabolism anomaly syndrome
93945	X-linked intellectual disability, Porteous type	85278	X-linked intellectual disability-craniofacial dysmorphism-epilepsy-ophthalmoplegia-cerebellar atrophy syndrome	163956	X-linked intellectual disability-nail dystrophy-seizures syndrome
→776	X-linked intellectual disability, Raymond type	163979	X-linked intellectual disability-craniofacioskeletal syndrome	2898	X-linked intellectual disability-plagiocephaly syndrome
3242	X-linked intellectual disability, Renpenning type	85280	X-linked intellectual disability-cubitus valgus-dysmorphism syndrome	85318	X-linked intellectual disability-precocious puberty-obesity syndrome
85285	X-linked intellectual disability, Schimke type	1568	X-linked intellectual disability-Dandy-Walker malformation-basal ganglia disease-seizures syndrome	3077	X-linked intellectual disability-psychosis-macroorchidism syndrome
3062	X-linked intellectual disability, Schutz type	2958	X-linked intellectual disability-dysmorphism-cerebral atrophy syndrome	85332	X-linked intellectual disability-retinitis pigmentosa syndrome
85323	X-linked intellectual disability, Seemanova type	94083	X-linked intellectual disability-dystonia-dysarthria syndrome		
85286	X-linked intellectual disability, Shashi type	85319	X-linked intellectual disability-epilepsy-progressive joint contractures-dysmorphism syndrome		
85324	X-linked intellectual disability, Shrimpton type				
85287	X-linked intellectual disability, Siderius type				
3063	X-linked intellectual disability, Snyder type				
85278	X-linked intellectual disability, South African type				
85325	X-linked intellectual disability, Stevenson type				
85288	X-linked intellectual disability, Stocco Dos Santos type				

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
3052	X-linked intellectual disability-seizures-psoriasis syndrome	383	X-linked mixed conductive and neurosensory hearing loss	85453	X-linked reticulate pigmentary disorder
457240	X-linked intellectual disability-short stature-overweight syndrome	383	X-linked mixed conductive and sensorineural deafness	1852	X-linked retinal dysplasia
→702	X-linked intellectual disability-spastic paraplegia with iron deposits syndrome	383	X-linked mixed conductive and sensorineural hearing loss	792	X-linked retinoschisis
163982	X-linked intellectual disability-spastic quadriplegia syndrome	383	X-linked mixed deafness with perilymphatic gusher	431272	X-linked scapuloperoneal muscular dystrophy
231692	X-linked isolated growth hormone deficiency	319605	X-linked MSMD	431272	X-linked scapuloperoneal syndrome
90625	X-linked isolated neurosensory deafness type DFN	319623	X-linked MSMD due to CYBB deficiency	86788	X-linked severe congenital neutropenia
90625	X-linked isolated neurosensory hearing loss type DFN	319612	X-linked MSMD due to IKBKG deficiency	75563	X-linked sideroblastic anemia
90625	X-linked isolated sensorineural deafness type DFN	319612	X-linked MSMD due to NEMO deficiency	2802	X-linked sideroblastic anemia and ataxia
90625	X-linked isolated sensorineural hearing loss type DFN	25980	X-linked myopathy with excessive autophagy	2802	X-linked sideroblastic anemia with ataxia
792	X-linked juvenile retinoschisis	178461	X-linked myopathy with postural muscle atrophy	99015	X-linked spastic paraplegia type 2
482606	X-linked keloid scarring-reduced joint mobility-increased optic cup-to-disc ratio syndrome	596	X-linked myotubular myopathy	100997	X-linked spastic paraplegia type 16
79447	X-linked lethal multiple pterygium syndrome	456328	X-linked myotubular myopathy-abnormal genitalia syndrome	171607	X-linked spastic paraplegia type 34
2148	X-linked lissencephaly type 1	85334	X-linked neurodegenerative syndrome, Bertini type	3175	X-linked spasticity-intellectual disability-epilepsy syndrome
452	X-linked lissencephaly with abnormal genitalia	85336	X-linked neurodegenerative syndrome, Hamel type	481	X-linked spinal and bulbar muscular atrophy
452	X-linked lissencephaly with ambiguous genitalia	314978	X-linked non progressive cerebellar ataxia	1145	X-linked spinal muscular atrophy type 2
452	X-linked lissencephaly-agenesis of the corpus callosum-genital anomalies syndrome	777	X-linked non-specific intellectual disability	404521	X-linked spinal muscular atrophy with respiratory distress
2442	X-linked lymphoproliferative disease	777	X-linked non-syndromic intellectual disability	85297	X-linked spinocerebellar ataxia type 3
1131	X-linked mandibulofacial dysostosis	90625	X-linked non-syndromic neurosensory deafness type DFN	85292	X-linked spinocerebellar ataxia type 4
1131	X-linked mandibulofacial dysostosis with limb anomalies	90625	X-linked non-syndromic neurosensory hearing loss type DFN	314978	X-linked spinocerebellar ataxia type 5
59306	X-linked McLeod syndrome	90625	X-linked non-syndromic sensorineural deafness type DFN	431272	X-linked SPMD
319605	X-linked mendelian susceptibility to mycobacterial diseases	293707	X-linked Ohdo syndrome	93349	X-linked spondyloepimetaphyseal dysplasia
319623	X-linked mendelian susceptibility to mycobacterial diseases due to CYBB deficiency	306597	X-linked Opitz BBB/G syndrome	168544	X-linked spondylometaphyseal dysplasia
319612	X-linked mendelian susceptibility to mycobacterial diseases due to IKBKG deficiency	306597	X-linked Opitz G/BBB syndrome	383	X-linked stapes gusher syndrome
319612	X-linked mendelian susceptibility to mycobacterial diseases due to NEMO deficiency	391330	X-linked Opitz syndrome	852	X-linked thrombocytopenia with normal platelets
435938	X-linked microcephaly-growth retardation-prognathism-cryptorchidism syndrome	363654	X-linked osteoporosis with fractures	3467	Xanthic urolithiasis
383	X-linked mixed conductive and neurosensory deafness	1175	X-linked parkinsonism-spasticity syndrome	93602	Xanthine dehydrogenase and xanthine aldehyde oxidase dual deficiency
		1652	X-linked progressive cerebellar ataxia	93601	Xanthine dehydrogenase deficiency
		83648	X-linked recessive hypercalciuric hypophosphatemic rickets	93601	Xanthine oxidase deficiency
		1652	X-linked recessive intellectual disability-macrocephaly-ciliary dysfunction syndrome	3467	Xanthine oxidoreductase deficiency
		54	X-linked recessive nephrolithiasis	93601	Xanthine stone disease
			X-linked recessive ocular albinism	93601	Xanthinuria type I
				93602	Xanthinuria type II
				158003	Xanthoma disseminatum
				79433	Xanthous oculocutaneous albinism
				79155	Xanthurenic aciduria
				67044	XDAT
				93602	XDH and AOX dual deficiency
				93601	XDH deficiency

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ORPHA number	Disease name	ORPHA number	Disease name	ORPHA number	Disease name
53351	XDP	243	XX-GD	3253	Zlotogora-Zilberman-Tenenbaum syndrome
293621	XECD	3375	XXX syndrome	913	Zollinger-Ellison syndrome
910	Xeroderma pigmentosum	168558	XY sex reversal-adrenal failure syndrome	2835	Zori-Stalker-Williams syndrome
90342	Xeroderma pigmentosum variant			912	ZS
→910	Xeroderma pigmentosum with neurologic manifestation	1770	XY type gonadal dysgenesis-associated anomalies syndrome	3474	Zunich-Kaye syndrome
220295	Xeroderma pigmentosum-Cockayne syndrome complex	2843	Xylitol dehydrogenase deficiency	295187	Zygodactyly type 1
75496	XGPT deficiency	75496	Xylosylprotein 4-beta-galactosyltransferase deficiency	295189	Zygodactyly type 2
181	XHED	370930	XYLT1-CDG	295191	Zygodactyly type 3
101088	XHIGM	2616	Yakut short stature syndrome	295193	Zygodactyly type 4
412069	Xia-Gibbs syndrome	99829	Yellow fever	295193	Zygodactyly, Castilla type
3469	XK aprosencephaly	99829	Yellow Jack	295189	Zygodactyly, Lueken type
452	XLAG (X-linked lissencephaly with abnormal genitalia) syndrome	662	Yellow nail syndrome	295191	Zygodactyly, Montagu type
		79434	Yellow oculocutaneous albinism	295187	Zygodactyly, Weidenreich type
596	XLCNM	3214	Yemenite deaf-blind hypopigmentation syndrome	73263	Zygomycosis
443197	XLDPP				
264580	XLG	707	Yersiniosis		
89936	XLH	99829	YF		
461	XLI	662	YNS		
596	XLMTM	876	Yolk sac tumor		
54	XLOA	252006	Yolk sac tumor of central nervous system		
306597	XLOS				
2442	XLP	252006	Yolk sac tumor of CNS		
443197	XLP	2828	YOPD		
85453	XLPDR	2255	Yorifuji-Okuno syndrome		
443197	XLPP	3240	Yoshimura-Takeshita syndrome		
792	XLRS	314485	Young adult-onset dHMN		
75563	XLSA	314485	Young adult-onset distal hereditary motor neuropathy		
2802	XLSA-A				
231393	XLTT	3471	Young syndrome		
25980	XMEA	3055	Young-Hughes syndrome		
317476	XMEN	2828	Young-onset Parkinson disease		
178461	XMPMA	477817	Yuan-Harel-Lupski syndrome		
93601	XO deficiency	3472	Yunis-Varon syndrome		
93601	XOR deficiency	319213	Zambian hemorrhagic fever		
910	XP	98912	ZASP-related myofibrillar myopathy		
220295	XP/CS complex	97240	Zebra body myopathy		
261476	Xp21 microdeletion syndrome	217017	Zechi-Ceide syndrome		
284180	Xp22.13p22.2 duplication syndrome	912	Zellweger syndrome		
1643	Xp22.3 microdeletion syndrome	369942	Zellweger-like contiguous gene deletion syndrome		
363654	XPDS				
90342	XPV	50812	Zellweger-like syndrome without peroxisomal anomalies		
314389	Xq12-q13.3 duplication syndrome	911	Zeta-associated-protein 70 deficiency		
1018	Xq22.3 microdeletion syndrome				
261483	Xq27.3-q28 microduplication syndrome	448237	Zika virus disease		
261483	Xq27.3q28 duplication syndrome	448237	Zika virus infection		
456328	Xq28 contiguous gene deletion syndrome	3301	Zimmer phocomelia		
		3473	Zimmermann-Laband syndrome		
243	XX female gonadal dysgenesis	439196	Zinc-responsive necrolytic acral erythema		
2855	XX gonadal dysgenesis-deafness syndrome	1775	Zinsser-Engman-Cole syndrome		
393	XX, male syndrome	3253	Zlotogora-Ogur syndrome		

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## List of diseases to be used instead of deprecated entities

→ Use these ORPHA number		instead of the deprecated entities	
ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
59	Allan-Herndon-Dudley syndrome	85337	X-linked intellectual disability, Zorick type
113	Bazex-Dupré-Christol syndrome	79458	Oley syndrome
113	Bazex-Dupré-Christol syndrome	79458	Congenital hypotrichosis-milia syndrome
138	CHARGE syndrome	1474	Colobomatous-microphthalmia-heart disease-hearing loss syndrome
138	CHARGE syndrome	1474	Hittner-Hirsch-Kreh syndrome
144	Lynch syndrome	99817	Non-polyposis Turcot syndrome
175	Cartilage-hair hypoplasia	1838	Metaphyseal dysplasia without hypotrichosis
175	Cartilage-hair hypoplasia	1838	Cartilage-hair hypoplasia-like-skeletal dysplasia without hypotrichosis syndrome
175	Cartilage-hair hypoplasia	93275	Thanatophoric dysplasia, Glasgow variant
193	Cohen syndrome	3084	Mirhosseini-Holmes-Walton syndrome
193	Cohen syndrome	3084	Pigmentary retinopathy-intellectual disability syndrome
193	Cohen syndrome	2829	Partington-Anderson syndrome
193	Cohen syndrome	3271	Radio-ulnar synostosis-retinal pigment abnormalities syndrome
193	Cohen syndrome	3271	Buntinx-Lormans-Martin syndrome
244	Primary ciliary dyskinesia	98861	Primary ciliary dyskinesia, Kartagener type
244	Primary ciliary dyskinesia	98861	Dextrocardia-bronchiectasis-sinusitis syndrome
244	Primary ciliary dyskinesia	98861	Immotile cilia syndrome, Kartagener type
244	Primary ciliary dyskinesia	98861	Kartagener syndrome
244	Primary ciliary dyskinesia	98861	Siewert syndrome
280	Wolf-Hirschhorn syndrome	85291	X-linked intellectual disability, Wittwer type
280	Wolf-Hirschhorn syndrome	85291	Wittwer syndrome
280	Wolf-Hirschhorn syndrome	98788	Pitt-Rogers-Danks syndrome
280	Wolf-Hirschhorn syndrome	98788	Intellectual disability-dysmorphism-intrauterine growth retardation syndrome
288	Hereditary elliptocytosis	98867	Hereditary pyropoikilocytosis
288	Hereditary elliptocytosis	98864	Common hereditary elliptocytosis
288	Hereditary elliptocytosis	98865	Homozygous hereditary elliptocytosis

→ Use these ORPHA number		instead of the deprecated entities	
ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
300	Bifunctional enzyme deficiency	2981	Pseudo-Zellweger syndrome
300	Bifunctional enzyme deficiency	2981	Thiolase deficiency
528	Berardinelli-Seip congenital lipodystrophy	1060	Systemic cystic angiomatosis-Seip syndrome
528	Berardinelli-Seip congenital lipodystrophy	1060	Brunzell syndrome
636	Neurofibromatosis type 1	3444	Watson syndrome
636	Neurofibromatosis type 1	3444	Pulmonic stenosis with 'café-au-lait' spots
636	Neurofibromatosis type 1	2029	Multiple non-ossifying fibromatosis
636	Neurofibromatosis type 1	2029	Jaffe-Campanacci syndrome
646	Niemann-Pick disease type C	79289	Niemann-Pick disease type D
646	Niemann-Pick disease type C	79289	Niemann-Pick disease, Nova Scotia type
672	Pallister-Hall syndrome	2113	Congenital hypothalamic hamartoma syndrome
672	Pallister-Hall syndrome	2113	CHHS
682	Hyperkalemic periodic paralysis	680	Normokalemic periodic paralysis
682	Hyperkalemic periodic paralysis	680	NormoKPP
682	Hyperkalemic periodic paralysis	680	NormoPP
682	Hyperkalemic periodic paralysis	680	Normokalemic PP
682	Hyperkalemic periodic paralysis	680	Periodic paralysis type 3
682	Hyperkalemic periodic paralysis	680	Potassium-sensitive normokalemic periodic paralysis
702	Pelizaeus-Merzbacher disease	85333	X-linked intellectual disability-spastic paraplegia with iron deposits syndrome
702	Pelizaeus-Merzbacher disease	85333	Arena syndrome
776	X-linked intellectual disability with marfanoid habitus	163953	X-linked intellectual disability, Raymond type
782	Axenfeld-Rieger syndrome	1831	De Hauwere syndrome
782	Axenfeld-Rieger syndrome	1831	De Hauwere-Chitty syndrome
782	Axenfeld-Rieger syndrome	1831	Iris dysplasia-hypertelorism-deafness syndrome
794	Saethre-Chotzen syndrome	1219	Aurocephalosyndactyly
794	Saethre-Chotzen syndrome	1219	Auralcephalosyndactyly
794	Saethre-Chotzen syndrome	1219	Kurczynski-Casperson syndrome
794	Saethre-Chotzen syndrome	3106	Robinow-Sorauf syndrome
798	Schinzel-Giedion syndrome	3118	Rudiger syndrome
823	Isolated spina bifida	93968	Meningocele
869	Triple A syndrome	99777	Achalasia-alacrimia syndrome
897	Waardenburg-Shah syndrome	918	ABCD syndrome

→ Use these ORPHA number		instead of the deprecated entities	
ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
897	Waardenburg-Shah syndrome	918	Albinism-black lock-cell migration disorder of the neurocytes of the gut-sensorineural deafness syndrome
910	Xeroderma pigmentosum	1569	De Sanctis-Cacchione syndrome
910	Xeroderma pigmentosum	1569	Xeroderma pigmentosum with neurologic manifestation
912	Zellweger syndrome	1271	Bowen syndrome
955	Acroosteolysis dominant type	2853	Serpentine fibula-polycystic kidneys syndrome
955	Acroosteolysis dominant type	2853	Exner syndrome
969	Acromicric dysplasia	2569	Moore-Federman syndrome
969	Acromicric dysplasia	2569	Dwarfism-stiff joint-ocular abnormalities syndrome
994	Fetal akinesia deformation sequence	995	X-linked fetal akinesia syndrome
994	Fetal akinesia deformation sequence	995	Holmes-Benacerraf syndrome
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	1401	CHAND syndrome
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	1401	Baughman syndrome
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	1401	CHANDS
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	1401	Curly hair-ankyloblepharon-nail dysplasia syndrome
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	3022	Rapp-Hodgkin syndrome
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	3022	Anhidrotic ectodermic dysplasia-cleft lip/palate syndrome
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	3022	Ectodermal dysplasia syndrome, Rapp-Hodgkin type
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	3022	Ectodermal dysplasia, Rapp-Hodgkin type
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	3022	RHS
1071	Ankyloblepharon-ectodermal defects-cleft lip/palate syndrome	99694	Alveolar synechia-ankyloblepharon-ectodermal dysplasia syndrome
1159	Progressive pseudorheumatoid arthropathy of childhood	2654	Syndesmodysplastic dwarfism
1159	Progressive pseudorheumatoid arthropathy of childhood	2654	Laplane-Fontaine-Lagardere syndrome

→ Use these ORPHA number		instead of the deprecated entities	
ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
1200	Choanal atresia-hearing loss-cardiac defects-craniofacial dysmorphism syndrome	77302	Oculo-oto-facial dysplasia
1215	Autosomal dominant optic atrophy plus syndrome	3349	Treft-Sanborn-Carey syndrome
1215	Autosomal dominant optic atrophy plus syndrome	3349	Optic atrophy-ophthalmoplegia-ptosis-deafness-myopathy syndrome
1215	Autosomal dominant optic atrophy plus syndrome	3212	Autosomal dominant optic atrophy and congenital deafness
1215	Autosomal dominant optic atrophy plus syndrome	3212	Konigsmark-Knox-Hussels syndrome
1234	Bartsocas-Papas syndrome	79446	Multiple pterygium syndrome, Aslan type
1263	Boomerang dysplasia	156723	Piepkorn dysplasia
1263	Boomerang dysplasia	156723	Short ribs-craniosynostosis-polysyndactyly syndrome
1299	Branchioskeletogenital syndrome	157788	Hypospadias-hypertelorism-coloboma and deafness syndrome
1359	Carney complex	623	NAME syndrome
1359	Carney complex	623	Nevi-atrial myxoma-myxoid neurofibromata-ephelides syndrome
1394	Cerebrofaciothoracic dysplasia	228407	Craniofacial dysmorphism-skeletal anomalies-intellectual disability syndrome
1394	Cerebrofaciothoracic dysplasia	228407	TMCO1 defect syndrome
1466	COFS syndrome	1317	CAMFAK syndrome
1466	COFS syndrome	1317	CAMAK syndrome
1466	COFS syndrome	1317	Cataract-microcephaly-arthrogryposis-kyphosis syndrome
1466	COFS syndrome	1317	Cataract-microcephaly-failure to thrive-kyphoscoliosis syndrome
1487	Cooks syndrome	2355	Kumar-Levick syndrome
1487	Cooks syndrome	2355	Nail dysplasia-camptodactyly-brachydactyly type B syndrome
1509	Coxopodopatellar syndrome	3112	Patella aplasia-coxa vara-tarsal synostosis syndrome
1643	Xp22.3 microdeletion syndrome	431	Ichthyosis-male hypogonadism syndrome
1658	Absence of fingerprints-congenital milia syndrome	1235	Ectodermal dysplasia-absent dermatoglyphs syndrome
1658	Absence of fingerprints-congenital milia syndrome	1235	Basan syndrome
1762	Trisomy Xq28	85281	MECP2 duplication syndrome
1762	Trisomy Xq28	85281	Lubs-Arena syndrome
1762	Trisomy Xq28	85281	X-linked intellectual disability, Lubs type
1762	Trisomy Xq28	85281	X-linked intellectual disability-hypotonia-recurrent infections syndrome

→ Use these ORPHA number		instead of the deprecated entities	
ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
1768	Familial caudal dysgenesis	1850	Renal dysplasia-megalocystis-sirenomelia syndrome
1768	Familial caudal dysgenesis	1850	Selig-Benacerraf-Greene syndrome
1855	Spondyloenchondrodysplasia	50816	Spondylometaphyseal dysplasia with combined immunodeficiency
1855	Spondyloenchondrodysplasia	50816	Roifman-Melamed syndrome
1855	Spondyloenchondrodysplasia	50816	SPENCDI
1855	Spondyloenchondrodysplasia	50816	Spondyloenchondrodysplasia with immune dysregulation
1896	EEC syndrome	1888	Ectrodactyly-ectodermal dysplasia without clefting syndrome
1896	EEC syndrome	1888	EEC syndrome without cleft lip/palate
1896	EEC syndrome	1889	Ectrodactyly-cleft palate syndrome
1896	EEC syndrome	1889	ECP syndrome
1896	EEC syndrome	2389	Lewis-Pashayan syndrome
1896	EEC syndrome	2389	Cleft lip/palate-ectrodactyly syndrome
1900	Ehlers-Danlos syndrome, kyphoscoliotic type	2691	Nevo syndrome
1900	Ehlers-Danlos syndrome, kyphoscoliotic type	2691	Cerebral gigantism, Nevo type
2036	Scalp-ear-nipple syndrome	3391	Odonto-onycho-hypohidrotic dysplasia-midline scalp defects syndrome
2036	Scalp-ear-nipple syndrome	3391	Ectodermal dysplasia-adrenal cyst syndrome
2036	Scalp-ear-nipple syndrome	3391	Tuffli-Laxova syndrome
2052	Fraser syndrome	2051	Fraser-like syndrome
2199	Epidermolytic palmoplantar keratoderma	496	Thost-Unna palmoplantar keratoderma
2199	Epidermolytic palmoplantar keratoderma	496	Non-epidermolytic palmoplantar keratoderma
2199	Epidermolytic palmoplantar keratoderma	89833	Palmoplantar keratoderma with tonotubular keratin
2353	Schilbach-Rott syndrome	1251	Blepharofacioskeletal syndrome
2353	Schilbach-Rott syndrome	1251	Richieri Costa-Guion Almeida-Rodini syndrome
2470	Matthew-Wood syndrome	91129	Anophthalmia-heart and pulmonary anomalies-intellectual disability syndrome
2510	Micro syndrome	2895	Pinsky-Di George-Harley syndrome
2510	Micro syndrome	2895	Microphthalmia-intellectual disability syndrome
2526	Microcephaly-lymphedema-chorioretinopathy syndrome	1432	Autosomal dominant chorioretinopathy-microcephaly syndrome
2609	Isolated complex I deficiency	936	Succinic acidemia
2616	3M syndrome	2661	Dwarfism-tall vertebrae syndrome

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ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
2637	Microcephalic osteodysplastic primordial dwarfism type II	46658	Primordial short stature-microdontia-opalescent and rootless teeth syndrome
2686	Cyclic neutropenia	2689	Intermittent neutropenia
2697	Arthrogryposis-renal dysfunction-cholestasis syndrome	1981	Fanconi syndrome-ichthyosis-dysmorphism syndrome
2697	Arthrogryposis-renal dysfunction-cholestasis syndrome	1981	Deal-Barrat-Dillon syndrome
2697	Arthrogryposis-renal dysfunction-cholestasis syndrome	3438	Biliary tract malformation-renal failure syndrome
2697	Arthrogryposis-renal dysfunction-cholestasis syndrome	3438	Cholestatic jaundice-renal tubular insufficiency syndrome
2697	Arthrogryposis-renal dysfunction-cholestasis syndrome	3438	Lutz-Richner-Landolt syndrome
2712	Oculofaciocardiodental syndrome	3013	Radiculomegaly of canine teeth- congenital cataract
2712	Oculofaciocardiodental syndrome	3013	Marashi-Gorlin syndrome
2750	Orofaciodigital syndrome type 1	90649	Orofaciodigital syndrome type 7
2750	Orofaciodigital syndrome type 1	90649	OFD7
2750	Orofaciodigital syndrome type 1	90649	Oral-facial-digital syndrome type 7
2750	Orofaciodigital syndrome type 1	90649	Whelan syndrome
2796	Pachydermoperiostosis	964	Acromegaly-cutis verticis gyrata-corneal leukoma syndrome
2882	Sitosterolemia	101022	Mediterranean macrothrombocytopenia
2909	Rothmund-Thomson syndrome	3333	Connective tissue dysplasia, Spellacy type
2909	Rothmund-Thomson syndrome	3333	Spellacy-Gibbs-Watts syndrome
2995	Baraitser-Winter cerebrofrontofacial syndrome	2649	Short stature-intellectual disability-eye anomalies-cleft lip/palate syndrome
2995	Baraitser-Winter cerebrofrontofacial syndrome	2649	Richieri Costa-Guion Almeida syndrome
2995	Baraitser-Winter cerebrofrontofacial syndrome	94084	Pachygyria-epilepsy-intellectual disability-dysmorphism syndrome
2995	Baraitser-Winter cerebrofrontofacial syndrome	94084	Cerebro-oculo-facial-lymphatic syndrome
2995	Baraitser-Winter cerebrofrontofacial syndrome	94084	Fryns-Aftimos syndrome
3057	Monoamine oxidase A deficiency	3065	X-linked intellectual disability-monoamine oxidase A metabolism anomaly syndrome



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ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
3157	Septo-optic dysplasia spectrum	1102	Anophthalmia-hypothalamo-pituitary insufficiency syndrome
3157	Septo-optic dysplasia spectrum	1102	14q22 microdeletion syndrome
3157	Septo-optic dysplasia spectrum	1102	Al Frayh-Facharzi-Haque syndrome
3157	Septo-optic dysplasia spectrum	1102	Monosomy 14q22
3157	Septo-optic dysplasia spectrum	1678	Dincsoy-Salih-Patel syndrome
3157	Septo-optic dysplasia spectrum	1678	Facial dysmorphism-ambiguous genitalia-hypopituitarism-short limbs syndrome
3157	Septo-optic dysplasia spectrum	2245	Hypopituitarism-postaxial polydactyly syndrome
3157	Septo-optic dysplasia spectrum	2245	Culler-Jones syndrome
3157	Septo-optic dysplasia spectrum	2243	Hypopituitarism-micropenis-cleft lip/palate syndrome
3157	Septo-optic dysplasia spectrum	2244	Hypopituitarism-microphthalmia syndrome
3157	Septo-optic dysplasia spectrum	2244	Kaplowitz-Bodurtha syndrome
3157	Septo-optic dysplasia spectrum	370006	Hypothalamic insufficiency-secondary microcephaly-visual impairment-urinary anomalies syndrome
3157	Septo-optic dysplasia spectrum	93943	Corpus callosum dysgenesis-hypopituitarism syndrome
3202	Dehydrated hereditary stomatocytosis	100039	Familial pseudohyperkalemia type 1
3253	Zlotogora-Ogur syndrome	90338	Margarita island ectodermal dysplasia
3447	Weaver syndrome	3446	Weaver-like syndrome
3460	Torg-Winchester syndrome	2775	Autosomal recessive carpotarsal osteolysis
3460	Torg-Winchester syndrome	2775	Hereditary multicentric osteolysis
3464	Woodhouse-Sakati syndrome	1011	Alopecia-hypogonadism-extrapyramidal syndrome
3464	Woodhouse-Sakati syndrome	1011	Devriendt-Legius-Fryns syndrome
3471	Young syndrome	1301	Bronchiectasis-oligospermia syndrome
33001	Lymphedema-distichiasis syndrome	1683	Distichiasis-congenital heart defects-peripheral vascular anomalies syndrome
33001	Lymphedema-distichiasis syndrome	2419	Lymphedema-ptosis syndrome
33364	Trichothiodystrophy	1245	BIDS syndrome
33364	Trichothiodystrophy	1245	Amish brittle hair syndrome
33364	Trichothiodystrophy	1245	Trichothiodystrophy type D
33364	Trichothiodystrophy	670	PIBIDS syndrome
33364	Trichothiodystrophy	670	Trichothiodystrophy type F
33364	Trichothiodystrophy	670	Trichothiodystrophy-sun sensitivity syndrome

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ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
33364	Trichothiodystrophy	453	IBIDS syndrome
33364	Trichothiodystrophy	453	Tay syndrome
33364	Trichothiodystrophy	453	Trichothiodystrophy type E
33364	Trichothiodystrophy	453	Trichothiodystrophy with congenital ichthyosis
33364	Trichothiodystrophy	2739	Onycho-tricho-dysplasia-neutropenia syndrome
33364	Trichothiodystrophy	2739	Itin syndrome
33364	Trichothiodystrophy	2739	ONMR syndrome
33364	Trichothiodystrophy	2739	Trichothiodystrophy type G
33364	Trichothiodystrophy	3123	Brittle hair syndrome, Sabinas type
33364	Trichothiodystrophy	3123	Brittle hair-mental deficiency syndrome
33364	Trichothiodystrophy	3123	Trichothiodystrophy type B
33364	Trichothiodystrophy	231256	Beta-thalassemia-trichothiodystrophy syndrome
33364	Trichothiodystrophy	75790	Pollitt syndrome
33364	Trichothiodystrophy	75790	Trichorrhexis nodosa syndrome
33364	Trichothiodystrophy	75790	Trichothiodystrophy type C
33364	Trichothiodystrophy	75790	Trichothiodystrophy-neurocutaneous syndrome
33364	Trichothiodystrophy	75789	SIBIDS syndrome
33364	Trichothiodystrophy	75789	Trichothiodystrophy-osteosclerosis syndrome
35069	Infantile neuroaxonal dystrophy	2174	Hunter-Carpenter-McDonald syndrome
36899	Myoclonus-dystonia syndrome	210566	Myoclonic dystonia 15
36899	Myoclonus-dystonia syndrome	210566	DYT15
36899	Myoclonus-dystonia syndrome	210566	Myoclonus-dystonia type 15
42738	Severe congenital neutropenia	37629	Neonatal neutropenia
42775	PHACE syndrome	1564	Dandy-Walker malformation-facial hemangioma syndrome
42775	PHACE syndrome	3195	Sternal malformation-vascular dysplasia syndrome
52368	Mohr-Tranebjaerg syndrome	3213	Deafness-opticoacoustic nerve atrophy-dementia syndrome
52368	Mohr-Tranebjaerg syndrome	3213	Jensen syndrome
52430	Inclusion body myopathy with Paget disease of bone and frontotemporal dementia	54238	Myotonic dystrophy type 3
56304	Atelosteogenesis type II	2640	Lethal short-limb dwarfism, McAlister-Crane type
56304	Atelosteogenesis type II	2640	McAlister-Crane syndrome
60030	Loeys-Dietz syndrome	97295	Furlong syndrome
60030	Loeys-Dietz syndrome	97295	Marfanoid habitus-craniosynostosis syndrome

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ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
69061	Idiopathic steroid-sensitive nephrotic syndrome	97552	Steroid-sensitive nephrotic syndrome without renal biopsy
69735	Hypotrichosis-lymphedema-telangiectasia-renal defect syndrome	2087	Glomerulonephritis-sparse hair-telangiectasis syndrome
79189	Peroxisome biogenesis disorder	34	Pipecolic acidemia
79189	Peroxisome biogenesis disorder	34	Hyperpipecolatemia
79259	Glycogen storage disease due to glucose-6-phosphatase deficiency type 1b	79261	Glycogen storage disease type 1D
79259	Glycogen storage disease due to glucose-6-phosphatase deficiency type 1b	79261	Type 1D glycogenosis
79259	Glycogen storage disease due to glucose-6-phosphatase deficiency type 1b	79260	Glycogen storage disease type 1C
79259	Glycogen storage disease due to glucose-6-phosphatase deficiency type 1b	79260	Type 1C glycogenosis
79452	Milroy disease	79450	Non-hereditary congenital primary lymphedema
79452	Milroy disease	79450	Milroy-like disease
79500	DOORS syndrome	1674	Digitorenocerebral syndrome
79500	DOORS syndrome	1674	DRC syndrome
79500	DOORS syndrome	1674	Eronen-Somer-Gustafsson syndrome
79502	Punctate palmoplantar keratoderma type 2	736	Palmoplantar porokeratosis of Mantoux
83628	PELVIS syndrome	2125	Sacral hemangiomas-multiple congenital abnormalities syndrome
85199	Craniosynostosis-anal anomalies-porokeratosis syndrome	2060	Fukuda-Miyanomae-Nakata syndrome
85293	X-linked intellectual disability, Cabezas type	85289	X-linked intellectual disability, Vitale type
86872	T-cell large granular lymphocyte leukemia	2687	Neutropenia-hyperlymphocytosis with large granular lymphocytes syndrome
90186	Meige disease	90185	Non-hereditary late-onset primary lymphedema
90186	Meige disease	90185	Meige-like disease
90340	Blaug syndrome	90341	Early-onset sarcoidosis
91387	Familial thoracic aortic aneurysm and aortic dissection	88636	Aortic dilatation-joint hypermobility-arterial tortuosity syndrome
93284	Spondyloepiphyseal dysplasia tarda	163673	Spondyloepiphyseal dysplasia, Byers type
93284	Spondyloepiphyseal dysplasia tarda	163673	Spondyloepiphyseal dysplasia-punctate corneal dystrophy syndrome

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ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
93950	X-linked intellectual disability, Sutherland-Haan type	93944	X-linked intellectual disability, Fichera type
95699	Congenital adrenal hyperplasia due to cytochrome P450 oxidoreductase deficiency	63269	Antley-Bixler syndrome with genital anomaly and disorder of steroidogenesis
95699	Congenital adrenal hyperplasia due to cytochrome P450 oxidoreductase deficiency	63269	Ambiguous genitalia-disordered steroidogenesis Antley-Bixler-like syndrome
95699	Congenital adrenal hyperplasia due to cytochrome P450 oxidoreductase deficiency	63269	Antley-Bixler syndrome type 2
95699	Congenital adrenal hyperplasia due to cytochrome P450 oxidoreductase deficiency	63269	Antley-Bixler syndrome, POR-related
97229	Riboflavin transporter deficiency	56965	Progressive bulbar paralysis of childhood
97229	Riboflavin transporter deficiency	56965	Fazio-Londe disease
97229	Riboflavin transporter deficiency	56965	Progressive bulbar palsy of childhood
98769	Spinocerebellar ataxia type 15/16	98770	Spinocerebellar ataxia type 16
98769	Spinocerebellar ataxia type 15/16	98770	SCA16
98772	Spinocerebellar ataxia type 19/22	101107	Spinocerebellar ataxia type 22
98772	Spinocerebellar ataxia type 19/22	101107	SCA22
98784	Autosomal dominant nocturnal frontal lobe epilepsy	98812	Paroxysmal hypnogenic dyskinesia
98784	Autosomal dominant nocturnal frontal lobe epilepsy	98812	Nocturnal paroxysmal dystonia
98784	Autosomal dominant nocturnal frontal lobe epilepsy	98812	Paroxysmal hypnagogic dyskinesia
98784	Autosomal dominant nocturnal frontal lobe epilepsy	98812	Paroxysmal hypnagogic dystonia
98784	Autosomal dominant nocturnal frontal lobe epilepsy	98812	Paroxysmal nocturnal dyskinesia
98808	Autosomal dominant dopa-responsive dystonia	101151	Dystonia 14
98808	Autosomal dominant dopa-responsive dystonia	101151	DYT14
98967	Schnyder corneal dystrophy	98968	Central discoid corneal dystrophy
168569	H syndrome	254723	Pigmented hypertrichosis with insulin-dependent diabetes mellitus syndrome
168569	H syndrome	254723	PHID

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ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
168569	H syndrome	254712	Familial sinus histiocytosis with massive lymphadenopathy
168569	H syndrome	254712	Familial Rosaï-Dorfman disease
168569	H syndrome	254712	Familial SHML
168569	H syndrome	254707	Faisalabad histiocytosis
168569	H syndrome	254707	FHC
182050	MYH9-related disease	850	May-Hegglin thrombocytopenia
182050	MYH9-related disease	850	MHA
182050	MYH9-related disease	850	May-Hegglin anomaly
182050	MYH9-related disease	850	May-Hegglin syndrome
182050	MYH9-related disease	1984	Fechtner syndrome
182050	MYH9-related disease	1984	Alport syndrome with leukocyte inclusions and macrothrombocytopenia
182050	MYH9-related disease	1019	Epstein syndrome
182050	MYH9-related disease	1019	Alport syndrome with macrothrombocytopenia
182050	MYH9-related disease	807	Sebastian syndrome
182050	MYH9-related disease	807	Macrothrombocytopenia with leukocyte inclusions
216866	Classic pantothenate kinase-associated neurodegeneration	157855	HARP syndrome
216866	Classic pantothenate kinase-associated neurodegeneration	157855	Hypoprebetalipoproteinemia-acanthocytosis-retinitis pigmentosa-pallidal degeneration syndrome
221061	Familial cerebral cavernous malformation	2486	Transverse limb deficiency-hemangioma syndrome
231568	Generalized dominant dystrophic epidermolysis bullosa	216989	Autosomal dominant dystrophic epidermolysis bullosa, Pasini type
231568	Generalized dominant dystrophic epidermolysis bullosa	216989	DDEB, Pasini type
231568	Generalized dominant dystrophic epidermolysis bullosa	79407	Autosomal dominant dystrophic epidermolysis bullosa, Cockayne-Touraine type
231568	Generalized dominant dystrophic epidermolysis bullosa	79407	DDEB, Cockayne-Touraine type
247691	Retinal vasculopathy and cerebral leukoencephalopathy	3421	Cerebroretinal vasculopathy
247691	Retinal vasculopathy and cerebral leukoencephalopathy	3421	CRV
247691	Retinal vasculopathy and cerebral leukoencephalopathy	3421	Grand-Kaine-Fulling syndrome
247691	Retinal vasculopathy and cerebral leukoencephalopathy	63261	HERNS syndrome

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ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
247691	Retinal vasculopathy and cerebral leukoencephalopathy	63261	Hereditary endotheliopathy-retinopathy-nephropathy-stroke syndrome
247691	Retinal vasculopathy and cerebral leukoencephalopathy	71291	Hereditary vascular retinopathy
247691	Retinal vasculopathy and cerebral leukoencephalopathy	71291	HVR
247691	Retinal vasculopathy and cerebral leukoencephalopathy	71291	Hereditary vascular retinopathy-Raynaud phenomenon-migraine syndrome
261483	Xq27.3q28 duplication syndrome	3423	Vasquez-Hurst-Sotos syndrome
261483	Xq27.3q28 duplication syndrome	3423	Hypogonadism-gynecomastia-X-linked intellectual disability syndrome
263463	CHST3-related skeletal dysplasia	1792	Humerospinal dysostosis
263463	CHST3-related skeletal dysplasia	93280	Spondyloepiphyseal dysplasia, Omani type
263463	CHST3-related skeletal dysplasia	93280	Humero-spinal dysostosis
264200	14q22q23 microdeletion syndrome	2055	Growth deficiency-brachydactyly-dysmorphism syndrome
264200	14q22q23 microdeletion syndrome	2055	Frias syndrome
284963	Marfan syndrome type 1	99715	MASS syndrome
284963	Marfan syndrome type 1	99715	Mitral valve-aorta-skeleton-skin syndrome
289825	Late-onset primary lymphedema	77242	Lymphedema tarda
289825	Late-onset primary lymphedema	77241	Lymphedema praecox
293843	3MC syndrome	2453	Malpuech syndrome
293843	3MC syndrome	2453	3MC3 syndrome
293843	3MC syndrome	2453	Malpuech facial clefting syndrome
293843	3MC syndrome	2506	Michels syndrome
293843	3MC syndrome	2506	3MC1 syndrome
293843	3MC syndrome	2506	Oculopalatoskeletal syndrome
293843	3MC syndrome	2998	Carnevale syndrome
293843	3MC syndrome	2998	3MC2 syndrome
293843	3MC syndrome	2998	Carnevale-Krajewska-Fischetto syndrome
293843	3MC syndrome	2998	Mingarelli syndrome
293843	3MC syndrome	2998	OSA syndrome
293843	3MC syndrome	2998	Oculo-skeletal-abdominal syndrome
293843	3MC syndrome	2998	Ptosis-strabismus-rectus abdominis diastasis syndrome
293864	Hypoplastic pancreas-intestinal atresia-hypoplastic gallbladder syndrome	137862	Martínez-Frías syndrome

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ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
293864	Hypoplastic pancreas-intestinal atresia-hypoplastic gallbladder syndrome	137862	Duodenal and extrahepatic biliary atresia-hypoplastic pancreas-intestinal malrotation syndrome
300751	Familial dilated cardiomyopathy with conduction defect due to LMNA mutation	83618	Severe dilated cardiomyopathy due to lamin A/C mutation
300751	Familial dilated cardiomyopathy with conduction defect due to LMNA mutation	83618	Severe dilated cardiomyopathy with or without myopathy
314632	Parkinsonism due to ATP13A2 deficiency	3336	Tomé-Brunet-Fardeau syndrome
319646	PGM1-CDG	711	Glycogen storage disease due to phosphoglucomutase deficiency
319646	PGM1-CDG	711	GSD due to phosphoglucomutase deficiency
319646	PGM1-CDG	711	GSD type 14
319646	PGM1-CDG	711	GSDXIV
319646	PGM1-CDG	711	Glycogen storage disease type 14
319646	PGM1-CDG	711	Glycogen storage disease type XIV
319646	PGM1-CDG	711	Glycogenosis due to phosphoglucomutase deficiency
319646	PGM1-CDG	711	Glycogenosis type 14
319646	PGM1-CDG	711	Glycogenosis type XIV
319646	PGM1-CDG	711	Phosphoglucomutase 1 deficiency
324737	SRD5A3-CDG	168972	Kahrizi syndrome
324737	SRD5A3-CDG	168972	Intellectual disability, Kahrizi type
324737	SRD5A3-CDG	168972	Intellectual disability-cataract-coloboma-kyphosis syndrome
324737	SRD5A3-CDG	139477	Al-Gazali-Dattani syndrome
329931	C3 glomerulonephritis	93559	C3 deposition glomerulonephritis without proliferation
331176	Autosomal recessive severe congenital neutropenia due to G6PC3 deficiency	178503	Dursun syndrome
331176	Autosomal recessive severe congenital neutropenia due to G6PC3 deficiency	178503	Pulmonary arterial hypertension-leukopenia-atrial septal defect syndrome
357225	Primary non-essential cutis verticis gyrata	1557	Cutis verticis gyrata-intellectual disability syndrome
357225	Primary non-essential cutis verticis gyrata	1557	McDowall syndrome
357225	Primary non-essential cutis verticis gyrata	217315	Cutis verticis gyrata-retinitis pigmentosa-sensorineural deafness syndrome

→ Use these ORPHA number		instead of the deprecated entities	
ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
357225	Primary non-essential cutis verticis gyrata	217315	Cutis verticis gyrata-retinitis pigmentosa-neurosensory deafness syndrome
357225	Primary non-essential cutis verticis gyrata	217315	Cutis verticis gyrata-retinitis pigmentosa-neurosensory hearing loss syndrome
357225	Primary non-essential cutis verticis gyrata	217315	Cutis verticis gyrata-retinitis pigmentosa-sensorineural hearing loss syndrome
357225	Primary non-essential cutis verticis gyrata	79482	Cutis verticis gyrata-thyroid aplasia-intellectual disability syndrome
357225	Primary non-essential cutis verticis gyrata	79482	Akesson syndrome
370114	Combined cervical dystonia	293838	Fatal infantile encephalopathy-pulmonary hypertension syndrome
370953	Congenital muscular dystrophy due to dystroglycanopathy	52428	Congenital muscular dystrophy type 1C
370953	Congenital muscular dystrophy due to dystroglycanopathy	52428	CMD1C
370953	Congenital muscular dystrophy due to dystroglycanopathy	52428	MDC1C
370953	Congenital muscular dystrophy due to dystroglycanopathy	98894	Congenital muscular dystrophy type 1D
370953	Congenital muscular dystrophy due to dystroglycanopathy	98894	MDC1D
399805	Male infertility with azoospermia or oligozoospermia due to single gene mutation	217034	Male infertility with normal virilization due to meiosis defect
399805	Male infertility with azoospermia or oligozoospermia due to single gene mutation	217034	Azoospermia due to maturation arrest
399805	Male infertility with azoospermia or oligozoospermia due to single gene mutation	217034	Azoospermia due to meiosis defect
399805	Male infertility with azoospermia or oligozoospermia due to single gene mutation	217034	Male infertility with normal virilization due to maturation arrest
399808	Male infertility with teratozoospermia due to single gene mutation	352613	Male infertility due to NANOS1 mutation
402041	Autosomal recessive distal renal tubular acidosis	93609	Autosomal recessive distal renal tubular acidosis without deafness
402041	Autosomal recessive distal renal tubular acidosis	93609	AR dRTA without deafness
402041	Autosomal recessive distal renal tubular acidosis	93609	AR dRTA without hearing loss



→ Use these ORPHA number		instead of the deprecated entities	
ORPHA number	Disease to be used	Deprecated ORPHA number	Deprecated entities
402041	Autosomal recessive distal renal tubular acidosis	93609	Autosomal recessive distal renal tubular acidosis without hearing loss
402041	Autosomal recessive distal renal tubular acidosis	93609	Distal renal tubular acidosis type 1c
402041	Autosomal recessive distal renal tubular acidosis	93609	dRTA type 1c
402041	Autosomal recessive distal renal tubular acidosis	93611	Autosomal recessive distal renal tubular acidosis with deafness
402041	Autosomal recessive distal renal tubular acidosis	93611	AR dRTA with deafness
402041	Autosomal recessive distal renal tubular acidosis	93611	AR dRTA with hearing loss
402041	Autosomal recessive distal renal tubular acidosis	93611	Autosomal recessive distal RTA with deafness
402041	Autosomal recessive distal renal tubular acidosis	93611	Autosomal recessive distal renal tubular acidosis with hearing loss
402041	Autosomal recessive distal renal tubular acidosis	93611	Distal renal tubular acidosis type 1b
402041	Autosomal recessive distal renal tubular acidosis	93611	dRTA type 1b
423693	Double outlet right ventricle with subaortic or doubly committed ventricular septal defect	99044	Double outlet right ventricle with subaortic ventricular septal defect
423693	Double outlet right ventricle with subaortic or doubly committed ventricular septal defect	99047	Double outlet right ventricle with doubly committed ventricular septal defect
444490	Familial chylomicronemia syndrome	411	Hyperlipoproteinemia type 1
448242	Autosomal recessive brachyolmia	93301	Brachyolmia type 1, Hobaek type
448242	Autosomal recessive brachyolmia	93303	Brachyolmia type 1, Toledo type
457059	Pseudohypoparathyroidism with Albright hereditary osteodystrophy	665	Albright hereditary osteodystrophy
457240	X-linked intellectual disability-short stature-overweight syndrome	3059	X-linked intellectual disability, Gu type
457240	X-linked intellectual disability-short stature-overweight syndrome	3059	MRX35

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