

מחלות נדירות

49 XXXXY

AADC – Aromatic Amino Acid Decarboxylase Deficiency

ABETALIPOPROTEINEMIA

ADA2 GENE MUTATION

ADAMS OLIVER SYNDROME

AIDS

AL KAISSI SYNDROME

ALAZAMI SYNDROME

ALBRIGHT HEREDITARY OSTEODYSTROPHY

ALDRICH SYNDROME

ALEXANDER SYNDROME

Allagile

ALLGROVE (TRIPLEA)

ALSTROM SYNDROME

Alstrom Syndrome

AMBIGUOUS GENITALIA

AMELOBLASTOMA

AMINO ACID DEHYDROGENASE (AADC) DEFICIENCY

APERT SYNDROME

AROMATIC L

AROMATIC L-AMINO DECARBOXYLASE DEFICIENCY - AADC DEF

Asphyxiating Thoracic Dystrophy (see also Jeune's)

AUTOSOMAL RECESSIVE CONGENITAL ICHTHYOSIS

BAINBRIDGE ROPERS SYNDROME

Barth Syndrome (3-Methyl Glutaconic Aciduria type II)

Bartter

BEALS SYNDROME

Bickel Fanconi Syndrome (Completely different from Fanconi anemia)

Blackfan Diamond Anemia

BORJESON-FORSSMAN-LEHMANN SYNDROME

Byler Dis.

C syndrome- Opitz Trigonoccephaly

CACNA1A MUTATION

CAMURATI ENGELMAN DISEASE

CAPILLARY MALFORMATION

CAPOS SYNDROME

Carbamoyl phosphate synthetase 1 deficiency

CARNITINE PALMITOYL TRASFERASE TYPE 2 DEF. SEVERE

INFANTILE FORM

CDH2 RELATED NEURODEVELOPMENTAL DISORDER

CDK10 SYNDROME

CDON DELETION

CEREBROTENDINOUS XANTOMATOSIS

Ch. 2 deletion 2 q12.2-2q13

CHARCOT MARIE TOOTH 2N-AARS MUTATION

CHILDHOOD APRAXIA OF SPEECH

CHOPS SYNDROME

Chronic Granulomatosis
CIPA
CITRULLINEMIA TYPE 2
CNOT3 MUTATION
COATS DISEASE
Cockayne
CODAS SYNDROME
COFFIN LOWRY SYNDROME
COFFIN SIRIS SYNDROME
COL11A1 GENE MUTATION
COL2A1 MUTATION
CONGENITAL CENTRAL HYPOVENTILATION SYNDROME
CONGENITAL CHLORID DIARRHEA CH. 7 MUTATION
Congenital dyserythropoiesis –see also Crigler Najar
Congenital insensitivity to pain with anhydrosis (CIPA)
CONGENITAL MYASTHENIA SYNDROME
CONGENITAL NON PROGRESSIVE CEREBELLAR ATAXIA
CONGENITAL SUCRASE
CRANIOMETAPHYSEAL DYSPLASIA
Crigler Najar -see also congenital dyserythropoiesis
CRMO
CTNNB1 MUTATION
CTSK GENE MUTATION
Cyclic Neutropenia
CYSTIC LYMPHANGIOMA
Cystinosis
DADA2
DASS
DEND
DHDDS MUTATION
DHFR DEF.
DIFFUSE PONTINE GLIOMA
DIGORGE SYNDROME
DIHIDROLIPOAMID DEHYDROGENASE DEFICIENCY
DNMT3A MUTATION
Donohue Syndrome (Leprechaunism)
Drash
DUE TO RYR1 MUTATION
DURAL SINUS MALFORMATION
Dyggve Melchior Clausen
DYSKERATOSIS CONGENITA
Dystonia Ziehen Oppenheim
ECHS1D
Ectopia cordis
Ectrodactyly ectodermal dysplasia clefting Syndrome (EEC)
EFTUD2 MUTATION SYNDROME
EHLER-DANLOS SYNDROME
Ehlers Danlos type IV

ENGELMANN 2 SYNDROME
Eosinophilic Fasciitis
EPIDERMOLYSIS BULLOSA
EPILEPTIC ENCEPHALOPATHY DUE TO ANTI FOLATE RECEPTOR
ANTIBODIES
EPP - ERYTHROPOETIC PROTOPORPHYRIA
ERYTHEMA ANNULARE
FACTOR 10 DEFICIENCY
FASCIO-SCAPULO-HUMERAL MUSCULAR DYSTROPHY
Fibrodysplasia Ossificans Progressiva
FIRES - FEBRILE INFECTION - RELATED EPILEPSY SYNDROME
Floating Harbor
FOXP1
Freeman Sheldon (see also Whistling face Syndrome)
FULL THICKNESS CORNEAL LACERATION
GASTROPARESIS
GENE ZNF292 MUTATION - DEVELOPMENTAL INTELLIGENCE DISORDER
GFER SYNDROME
GILLESPIE SYNDROME
GLANZMAN THROMBASTHENIA
GLANZMANN THROMBASTHENIA - LIKE
GLASS SYNDROME
GLOBAL DEVELOPMENTAL DELAY - CLTC RELATED
GLUT 1 DEFICIENCY
GLUTARIC ACIDURIA TYPE 1
Glutaric Aciduria type II
GLYCOGEN STORAGE DISEASE TYPE 9A
GNAS GENE MUTATION - PSEUDOHYPOPARATHYROIDISM
Gorham disease
GREIG CEPHALOPOLYSYNDACTYLY SYNDROME
GRIN2B MUTATION
H SYNDROME
H.I.V.
HELSMOOTEL VAN DER A SYNDROME
HEMIIHYPERPLASIA-LIPOMATOSIS SYNDROME
HEREDITARY FRUCTOSE INTOLERANCE -ALDOB GENE MUTATION
HEREDITARY PARAGANGLIOMA PHEOCHROMOCYTOMA SYNDROME
HEREDITARY SENSORY AUTONOMIC NEUROPATHY
HEREDITARY SPASTIC PARAPARESIS
HEREDITARY SPASTIC PARAPLEGIA
HERMANSKY PUDLAK SYNDROME 5 - HPS5
HETEROZYGOUS FOR ASXL3 MUTATION
HHH Syndrome
HIBCH SYNDROME
Histiocytosis X
Hunter
Hyper IgE (Job's)
HYPERCHOLANEMIA
Hypercholesterolemia Familial Homozygous

Hyperoxaluria type I
Ichthyosis:(specific types only) Lamellar, Non-bullous Congenital, Ichthyosis erythroderma
IDIOPATHIC ACUTE TRANSVERSE MYELITIS
IGF1R AND HNRNP2A GENE MUTATIONS
IMAGE SYNDROME
IMMUNODEFICIENCY 14A AUTOSOMAL DOMINANT
Incontinentia Pigmenti
Infantile Polyarthritis Nodosa
INTELLECTUAL DEVELOPMENTAL DISORDER WITH SEVERE SPEECH AND AMBULATION DEFECTS SYNDROME
INTRACTABLE DIARRHEA OF INFANCY SYNDROME
IPEX
ISOMALTASE DEFICIENCY
ISOVALERIC ACIDEMIA
Isovaleric Acidemia
ISOVALERIC ACIDURIA
Jacobsen Syndrome
Jeune's (Asphyxiating Thoracic Dystrophy)
Job's (Hyper IgE)
JOUBERT SYNDROME
KABUKI SYNDROME
KAT6A SYNDROME
KBG SYNDROME
KDM5C GENE MUTATION
Kearns Sayre
Kenny-Caffey Syndrome
KINDLER SYNDROME
KINSHIP SYNDROME
KLEEFSTR SYNDROME
KLEEFSTR2 SYNDROME
KLINE SYNDROME
KLIPPLE TRENAUNAY SYNDROME
KOHLER SYNDROME-COWDEN
Kostman Disease
KOSTMANN SYNDROME
Krabbe Disease
LADD Lacrimo-auricular-dento-digital Syndrome
LANDAU-KLEFFNER SYNDROME
Langerhans Cell Dis (see also Histiocytosis X)
Larsen Syndrome
LEGIUS SYNDROME
Leucocyte Adhesion Deficiency Type 2
Leucodystrophy Van der Knapp
LEUKOCYTE ADHESION DEFICIENCY TYPE 2
LIPOAMIDE DEHYDROGENASE DEFICIENCY (E3)
LIPOPROTEIN LIPASE DEFICIENCY
LOEYS-DIETZ SYNDROME
LOEYS-DIETZ SYNDROME

LONG-CHAIN 3HYDROXY-ACYL-COA DEHYDROGENASE DEFICIENCY-
LCHADD
LOWE SYNDROME
Lymphoma congenital
LYSINURICPROTEIN INTOLERANCE (LPI)
MACROCEPHALY CAPILLARY MALFORMATION
MAJEED SYNDROME
MALAN (SOTOS LIKE) SYNDROME
MARS MUTATION
MCCUNE ALBRIGHT SYNDROME
MCM SYNDROME - MACROCEPHALY-CAPILLARY MALFORMATION
MECOM - GENE MUTATION
MECP2 GENE MUTATION
MECR MUTATION
MEF2C HAPLOINSUFFICIENCY SYNDROME
MEN2B MUTATION
MEPAN SYNDROME
METHYL MALONIC ACIDURIA
MILROY DISEASE
MIXED GONADAL DYSGENESIS
MMP 9 SYNDROME
MOEBIUS SYNDROME
Morquio Syndrome
Moya-Moya Disease
MRFACD SDR
MSUD
MUCKLE WELLS SYNDROME
Mucopolysaccharidosis type II, type IVa, type IVb & type VI
MULTIPLE HEREDITARY EXOSTOSIS AND OSTEOCHONDROMATOSIS
MUSCLE EYE BRAIN DISEASE
MUTATION IN CUL4B GENE
MYCOSIS FUNGOIDES
MYH7 GENE MUTATION
MYHRE SYNDROME
Myhre Syndrome
NEONATAL PROGERIA
Nephrogenic Diabetes Insipidus (only X-linked N.D.I)
Netherton Syndrome
NEVUS UNIVUS LATERIS
NFIA MUTATION - BRAIN MALFORMATION WITH OR WITHOUT
URINARY TRACT MALFORMATION
NICOLAIDES-BARAITSER SYNDROME
NIEMANN PICK C
NOONAN SYNDROME 7- CARDIOFACIOCUTANEOUS SYNDROME
NSD1 MUTATION
O'DONNELL-LURIA-RODAN SYNDROME
OCCULO FACIAL DENTAL SYNDROME
OGDEN SYNDROME
OHDO SYNDROME SBBYS VARIANT

OLLIER DISEASE
Omenn
Pallister-Hall Syndrome
PAN - Periarthritis Nodosa
Papillorenal Syndrome (Pax2 gene mutation)
PARKES WEBER SYNDROME
Pearson marrow pancreas syndrome
PEDIATRIC MULTIPLE SCLEROSIS
Pelizaeus-Merzbacher Disease
PENTA X
PEPCK-C OMIM 26680
PERCC1
PERIODIC PARALYSIS - DUE TO RYR1 MUTATION
PFEIFFER SYNDROME
PHELAN MCDERMID SYNDROME
PHF21A GENE MUTATION
PHOSPHOGLUCOMUTASE 1 DEFICIENCY
PICNODYSOSTOSIS
PITT HOPKINS LIKE SYNDROME
PITT HOPKINS SYNDROME
PITUITARY STALK INTERRUPTION SYNDROME
PLP1 MUTATION
POLAND SYNDROME
POPLITEAL PTERYGIUM SYNDROME
POTOKI-LUPSKI SYNDROME
PPB - PLEUROPULMONARYBLASTOMA
PRIMARY HYPEROXALURIA TYPE 1
Progyria
PROLIDASE DEFICIENCY
PRSS1 MUTATION
Pseudo-hypo-aldosteronism
PSEUDOHYPOALDOSTERONISM TYPE 1B
PSEUDOHYPOPARATHYROIDISM TYPE2
PTEN MUTATION
PTLD (post transplantation lymphoproliferative dis)
PUDLAK TYPE 2 SYNDROME
PULMONARY ALVEOLAR PROTEINOSIS
PURA SYNDROME
Pycnodysostosis
Pyruvate Dehydrogenase deficiency
RENPENNING SYNDROME - PQBP1 MUTATION
RETICULATE PIGMENTED DISORDER
Rogers Syndrome
RT TMJ ANKYLOSIS
SATB2 ASSOCIATED SYNDROME
SCHAAF - YANG SYNDROME
Schimke Immuno-Osseous Dysplasia
SCHINZEL GIEDION

SCHUURS JMAKERS SYNDROME
SCIMITAR SYNDROME
SCN4A MUTATION
SCYL1 MUTATION - SPINOCEREBELLAR ATAXIA TYPE 21
SEPIAPTERIN REDUCTASE DEFICIENCY
SETD5 MUTATION
SHANK 2 MUTATION
SHANK3 GENE MUTATION
SHORT SYNDROME
SIALURIA MIM 269921
SIFRIM-HITS-WEISS SYNDROME
SIFRIM-HITZ-WEISS SYNDROME
SILVER RUSSELL SYNDROME
SLC9A7 GENE MUTATION
SLO -SMITH LEMLI OPITZ SYNDROME
SMARCA 2 MUTATION
SMITH KINGSMORE SYNDROME
SMOOTH MUSCLE DYSFUNCTION - ACTA2 MUTATION
SNIJDERS BLOCK CAMPEAU SYNDROME
SOX2 MUTATION
SPONDYLO METAPHYSEAL DYSPLASIA - VARIANT RPL 13
Spondyloepiphyseal Dysplasia
STAT 3 MUTATION
Stevens Johnson
SUCRASE ISOMALTASE DEFICIENCY
SYNGAP1- RELATED INTELLECTUAL DISABILITY
SYP1
SYSTEMIC MASTOCYTOSIS
Systemic Scleroderma
SZT2 GENE MUTATION
Takayasu Vasculitis
TAR SYNDROME
TATTON-BROWN-RAHMANSYNDROME - TBRS
TEC -PR2 SYNDROME - OBSTRUCTIVE SLEEP APNEA SYNDROME
TRICHOHEPATOENTERIC SYNDROME
TRICHORHINOPHALANGEAL SYNDROME
TRMT10A RELATED DISORDER
TUBER CINEREUM HAMARTOMA
TYROSINEMIA TYPE 2
TYROSINEMIA TYPE 3
Tyrosinemia type I
Tyrosinemia type II
Upshaw Schulman Syndrome
USP7 GENE MUTATION
Van der Knapp (see also Leucodystrophy)
VERHEIJ SYNDROME
VERVERI - BRADY SYNDROME
VICI SYNDROME

VON WILLEBRANDS DIS.
Wegener Granulomatosis
WEISS- KRUSZKA SYNDROME
Widemann Rautenstrauch Syndrome
WIEDEMANN-STEINER SYNDROME
Wiskott Aldrich
WISKOTTALDRICH SYNDROME
X LINKED HYPOPHOSPHATEMIC RICKETS
X LINKED INTELLECTUAL DISABILITY - SIDERIUS TYPE
Xeroderma Pigmentosum
XIA-GIBBS SYNDROME
Ziehen Oppenheim Dystonia
ZNF292
ZNF335 -GENE MUTATION
ZTTK
1.