

Established Conditions for the Purposes of CDS Early Intervention Eligibility

All children diagnosed with one of the following established conditions, which have a high probability of resulting in developmental delay, are eligible for early intervention services until their third birthday.  A child’s medical record may be used to establish eligibility, without conducting an evaluation of the child, if the record indicates that the child has an established condition found on the list below. Please note that this list is not exhaustive and other established conditions may be used for eligibility with reliable information from reputable sources.

3-Hydroxy-3-Methylglutaric Aciduria (HMG)

3-Methylcrotonyl-CoA Carboxylase Deficiency (3MCC)

Abusive Head Trauma (*previously known as “Shaken Baby Syndrome”*)

Achondrogenesis

Acrocallosal Syndrome

Acrodysotosis

Acute Disseminated Encephalomyelitis

Adrenoleukodystrophy

Agenesis of Corpus Callosum

Aicardi Syndrome

Aicardi-Goutieres Syndrome

AIDS

Albinism

Alexander Disease

Allan Herndon Syndrome

Alpers Syndrome

Alternating Hemiplegia

Amyelia

Amyoplasia Congenita

Anencephaly

Angelman Syndrome

Aniridia-Cerebellar Ataxia Syndrome

Anopthalmos/Micropthalmos

Anterior Horn Cell Disease

Antimongolism Syndrome

Apert Syndrome

Antley-Bixler Syndrome

APGAR <=3 @20 minutes

Aphasia

Apraxia

Argininemia (ARG)

Argininosuccinate Aciduria (ASA)

Arhinencephaly

Arnold-Chiari Syndrome Type II

Arthrogryposis

Ataxia

Ataxia-Telangiectasia Syndrome

Attachment Disorders

Attention Deficit Disorder (ADD)

Attention Deficit Hyperactivity Disorder (ADHD)

Autism Spectrum Disorder (ASD)

Anxiety Disorder

Baller Gerold Syndrome

Bannayan Riley Ruvalcaba

Bardet-Beidl Syndrome

Bartoscas-Papas Syndrome

Basal Cell Nevus Syndrome

Batten Disease

B-Ketothiolase Deficiency (BKT)

Biotinidase Deficiency (BIOT)

Blindness

Bloch-Sulzberger Syndrome

Bobble Head Doll Syndrome

Borjeson Syndrome

Brain Malformation

Brain Tumor

Brancio-Oto-Renal Syndrome

Bulbar Palsy

C Syndrome

Canavan Disease

Cancer

Carbohydrate Deficient Glycoprotein Syndrome

Carbohydrate Metabolism Disorder

Cardio-Facio-Cutaneo Syndrome

Carnitine Palmityl Transferase Deficiency Type II (CPTII)

Carnitine Uptake Defect/Carnitine Transport Defect (CUD)

Carpenter Syndrome

Caudal Regression Syndrome

Central Core Disease

Cerebellar Agenesis

Cerebral Ataxia

Cerebral Atrophy

Cerebral Degeneration

Cerebral Dysgenesis

Cerebral/Cortical Dysplasia

Cerebral Gigantism

Cerebral Palsy

Cerebral Venous Thrombosis

Cerebrocostomandibular Syndrome

Cerebrovascular accident

CHARGE Syndrome

Chiari Malformation

Childhood Disintegrative Disorder

Chromosomal Disorders

Chromosome 10 Duplication

Chromosome 10p+

Chromosome 11p-

Chromosome 12p-

Chromosome 13q-

Chromosome 13q+

Chromosome 14 Deletion

Chromosome 14 Ring

Chromosome 15 Ring

Chromosome 15, Distal Trisomy

Chromosome 15q11-q13

Chromosome 16 Duplication

Chromosome 17q12 Duplication

Chromosome 18 Ring

Chromosome 18, Tetrasomy 18p

Chromosome 18p-

Chromosome 18q-

Chromosome 19p Duplication

Chromosome 2p21

Chromosome 20q Trisomy

Chromosome 21

Chromosome 21q-

Chromosome 22 Ring

Chromosome 22, Trisomy Mosaic

Chromosome 22q-

Chromosome 22q11.2 Duplication

Chromosome 22q11.2 Deletion

Chromosome 1p36 Deletion

Chromosome 2q32 Deletion

Chromosome 3, Monosomy 3p2

Chromosome 3, Trisomy 3q2

Chromosome 3q+

Chromosome 4 Ring

Chromosome 4, Monosomy 4q

Chromosome 4, Monosomy Distal 4q

Chromosome 4, Partial Trisomy Distal 4q

Chromosome 4q-

Chromosome 4 q+

Chromosome 4, Trisomy 4p

Chromosome 5, Trisomy 5p

Chromosome 5 Deletion

Chromosome 5p-

Chromosome 6 Ring

Chromosome 6, Partial Trisomy 6q

Chromosome 7, Partial Monosomy 7p

Chromosome 7, Partial Duplication Syndrome

Chromosome 7, Partial Monosomy

Chromosome 8, Monosomy 8p2

Chromosome 9 Ring

Chromosome 9 Trisomy

Chromosome 9p Deletion

Chromosome 9 Tetrasomy 9p

Chromosome 9, Trisomy 9pter-q22-32

Chromosome Xq26.2 Duplication

Chromosome 49XXXXY Syndromes

Citrullinemia

Cleft Lip

Cleft Palate

Club Foot

CNS Degenerative Disorder

Cockayne Syndrome

Coffin-Lowry Syndrome

Coffin-Siris Syndrome

Congenital Bilateral Persylvian

Congenital Cataracts

Congenital Fiber Type Disproportion

Congenital Heart Disease

Congenital Heart Defect

Congenital Muscular Dystrophy

Congenital or Acquired Amputation of Limb

Conjoined Twin

Cornelia de Lange Syndrome

Costello Syndrome

Craniosynostosis

Cytomegalovirus Disease, Congenital (CMV)

Dandy Walker Syndrome

De Barsy Syndrome

De Sanctis Cacchione Syndrome

Deafness

Dejerine Sottas Disease

Depression of Infancy or Childhood

Deprivation Maltreatment Disorder

Diencephalic Syndrome

DiGeorge Syndrome

DOOR Syndrome

Down Syndrome

Dravet Syndrome

Dubowitz Syndrome

Duchenne Muscular Dystrophy

Dyggve-Melchior-Clausen Syndrome

Early Onset Torsion Dystonia

Emmanuel Syndrome

Encephalitis

Encephalocele

Encephalopathy, Neonatal Hypoxic Ischemic Encephalopathy

Encephalopathy, Static

Epidermal Nevus Syndrome

Epilepsy

Epstein’s Syndrome

Ethmocephaly

Fabry’s Disease

Fahr’s Disease

Failure to Thrive

Familial Retardation Syndrome

Farber Disease

Fatty Acid Storage Disease

Fazio-Londe Disease

Feeding Tube Dependent (Nasal or Abdominal)

Fetal Alcohol Spectrum Disorders (FASD)

Fetal Hydantoin Syndrome

Fetal Valproate Syndrome

FG Syndrome

Filippi Syndrome

Floating Harbor Syndrome

Fountain Syndrome

Fragile X Syndrome

Friedreich’s Ataxia

Fryns Syndrome

Fucosidosis

Galactosemia

Gangliosidosis

Gaucher Syndrome

Glutaric Acidemia Type I (GAI)

Glutaric Acidemia Type II (GAII)

Hallervoden-Spatz Disease

Hallgren Syndrome

Harlequin Fetus Syndrome

Hearing Impairment, significant/progressive

Hemimegalencephaly

Hemiparesis

Hemoglobin Disorders/ Sickle Cell Disease (Hb SS, Hb S/B, and Hb S/C)

Herpes Simplex, Congenital

HIV Positive

Holoprosencephaly

Homocystinuria (HCY)

Human HOXA 1 Syndromes

Hunter Syndrome

Hurlur-Sheie Disease

Hydroencephaly

Hydrocephalus

Hyperornithinemia Hyperammoninemia Homocitrullinemia Syndrome (HHH)

Hypoglossia Hypodactylia Spectrum

Hypoxic Ischemic Encephalopathy

Hypsarrhythmia

I Cell Disease

Inborn Errors of Metabolism

Incontinentia Pigmenti Syndrome

Infantile Anorexia

Infantile Neuroaxonal Dystrophy

Infantile Spasms

Iniencephaly

Intellectual Disability

Intracranial Hemorrhage

Intrauterine Growth Restriction (<=10th percentile for gestational age)

Intraventricular Hemorrhage

Ischemic Brain Infarction

Isovaleric Acidemia (IVA)

Jacobsen Syndrome

Jansky-Bielschowsky Disease

Jervell and Lange-Nielson Syndrome

Johanson-Blizzard Syndrome

Joubert Syndrome

Juberg-Marsidi Syndrome

Kabuki Syndrome

KBG Syndrome

Keratitis Ichthyosis Deafness Syndrome

Kernicterus

Kinsbourne Syndrome

Kleefstra Syndrome

Klinefelter Syndrome

Krabbe’s Disease

Kufs Disease

Kugelberg-Welander Syndrome

L 1 Syndrome

Lambert-Eaton Myasthenic Syndrome

Landau Kleffner Syndrome

Langer-Giedion Syndrome

Laurence Moon Syndrome

Lead in the Blood (*previously known as “Lead Poisoning”*)

Lead Poisoning (*now referred to as “Lead in the Blood”)*

Leber’s Congenital Amaurosis

Leigh’s Disease

Lennox-Gastaut Syndrome

Lenz Microphthalmia Syndrome

Leprechaunism

Lesch-Nyhan Syndrome

Leukodystrophy

Linear Sebaceous Nevus Syndrome

Lipid Metabolism Disorders

Lipodystrophy, Congenital

Lissencephaly Syndrome

Locked in Syndrome

Long Chain Fatty Acid Storage Disease

Long-Chain L-3 Hydroxyacyl-CoA Dehydrogenase Deficiency (LCHAD)

Low Birth Weight <1200 grams

Low Vision

Lowe Syndrome

Lysosomal Storage Disorders

Macrocephaly

Malignant Neoplasm of Brain

Mandibulofacial Dysotosis with Microcephaly

Mannosidosis

Maple Syrup Urine Disease (MSUD)

Marden Walker Syndrome

Marinesco Sjorgen Syndrome

Maroteaux-Lamy Syndrome

Marshall-Smith Syndrome

Maxillofacial Dysotosis

Meckel Disease

Medium-Chain Acyl-CoA Dehydrogenase Deficiency (MCAD)

MELAS Syndrome

Melnick Frazier Syndrome

Meningitis

Meningocele

Meningomyelocele

Menkes Syndrome

Mercury Toxicity

MERRF Syndrome

Methylmalonic Acidemia (MUT)

Methylmalonic Acidemia Cobalamin A, B (Cbl A, B)

MHBD Deficiency

Microcephaly

Midas Syndrome

Miller-Diecker Syndrome

Mitochondrial Disorder

Mobius Sequence

Mohr-Tranebjaerg Syndrome

Morquio Syndrome

Motor Neuron Disease

Mowat-Wilson Syndrome

MPPH Syndrome

Mucolipidosis

Multi System Developmental Disorder

Multiple Anomalies of Brain

Multiple Sulfatase Deficiency

Muscular Dystrophy

Myasthenia, Congenital

Myasthenia Gravis, Neonatal Transient

Myelocele

Myyelomeningocele

Myelocystocele

Myhre Syndrome

Mucopolysaccharidosis Type I

Myopathy, Congenital

Myopathy Storage Disease

Myositis Ossificans Progressive

Myotonic Dystrophy

Myotubular Myopathy

Nager-de-Reynier Syndrome

Nasal Hypoplasia MR/ID

Nemaline Rod Myopathy

Neonatal Abstinence Syndrome (NAS)

Neonatal Opioid Withdrawal Syndrome (NOWS)

Neu-Laxova Syndrome

Neural Tube Defect

Neurofibromatosis

Neuronal Ceroid-Lipofuscinoses-Amaurotic Syndrome

Neuropathy, Ataxia and Retinitis Pigmentosa

Neuropathy, Congenital Hypomyelination

Neuropathy, Giant Axonal

Neuropathy, Hereditary Sensory

Neuropathy, Peripheral

Niemann Pick

Non-Ketotic Hyperglycemia

Norrie’s Syndrome

Oculocerebral Syndrome with Hypopigmentation

Oculocerebrocutaneous Syndrome

Oculocerebellar Atrophy, Hereditary

Ohtahara Syndrome

Olivopontocerebellar Atrophy, Hereditary

Opitz G/BBB Syndrome

Opitz Syndrome

Oral-Facial Digital Syndrome

Organic Acidemias

Ornithine-Carbamyl Transferase Deficiency

Orotic Aciduria

Osteopetrosis

Otopalatodigital Syndrome

Pachgyria

Pallister W Syndrome

Pallister-Killian Syndrome

Paraplegia, Hereditary Spastic

Partial-Cerebellar Ataxia MR/ID

Pediatric AIDS/HIV

Pediatric Feeding Disorders

Pelizaeus-Merzbacher Disease

Pentasomy X Dysotosis

Periventricular Leukomalacia

Peroxisomal Disorders

Phelan-McDermid Syndrome

Phenylketonuria (PKU)

Phocomelia

Pick’s Disease

Poliomyelitis

Polymicrogyria

Pompe Disease

Porencephaly

Post-Traumatic Stress Disorder

Potocki-Lupski Syndrome

Prader-Willi Syndrome

Prematurity <29 weeks

Prenatal Alcohol Exposure (PAE)

Progressive Multi-focal Leukoencephalopathy

Progressive Myoclonic Epilepsy

Propionic Acidemia (PROP)

Pseudo-Hurler Polydystrophy

Pyruvate Carboxylase Deficiency

Pyruvate Dehydrogenase Deficiency

Quadriplegia

Reactive Attachment Disorder

Refsum Syndrome

Regulatory Disorders of Sensory Processing

Rett Syndrome

Rhizomelic Chondrodysplasia Punctata

Robert’s Syndrome

Rosenberg Chutorian Syndrome

Roussy Levy Syndrome

Rubella, Congenital

Rubenstein-Taybi Syndrome

Russel-Silver Syndrome

Sandhoff Disease

Sanfilippo Syndrome

Santavuori Disease

Scheie Syndrome

Schilder’s Disease

Schindler Disease

Schinzel Giedion Syndrome

Schizencephaly

Schwartz-Jampel Syndrome

Scott Craniodigital Syndrome

Seckel Syndrome

Semilobar Holoprosencephaly

Shaken Baby Syndrome (*now referred to as “Abusive Head Trauma”)*

Shprintzen-Goldberg Craniosynostosis Syndrome

Sialidosis

Sickle Cell Disease/Hemoglobin Disorders (Hb SS, Hb S/B, and Hb S/C)

Simpson Dysmorphia Syndrome

Sirenomelia Sequence

Sjogren-Larsson Syndrome

Sly Syndrome

Small for Gestational Age <=10th percentile

Smith-Lemli-Opitz Syndrome

Smith-Magenis Syndrome

Social Anxiety Disorder

Sotos Syndrome

Spastic Paraplegia

Sphingolipidoses

Spielmeyer-Vogt Disease

Spina Bifida

Spinal Cord Injuries

Spinal Lipomeningocele

Spinal Muscular Atrophy

Spinocerebellar Disorders

Steinert-Myotonic Dystrophy Syndrome

Stickler Syndrome

Stroke

Sturge-Weber Syndrome

Subacute Sclerosing Panencephalitis

Substance Exposed Infant (SEI)

Succinic Semialdehyde Dehydrogenase Deficiency

Sulfatide Lipidosis

Sydenham Chorea

Syphilis, Congenital

Syringohydromyelia

Tay-Sachs Disease

Tetralogy of Fallot

Timothy Syndrome

Torticollis

Toxoplasmosis, Congenital

Transposition of the Great Vessels

Transverse Myelitis

Traumatic Brain Injury

Trifunctional Protein Deficiency (TFP)

Triphosphate Isomerase Deficiency

Triple X Syndrome

Triploidy Syndrome

Trisomy 12p

Trisomy 13

Trisomy 18

Trisomy 3

Trisomy 8

Trisomy 9

Tuberous Sclerosis

Tyrosinemia Type I and Type II

Urea Cycle Defect

Usher Syndrome

VACTERL Syndrome with Hydrocephalus

Velocardiofacial Syndrome

Ventilator Dependent

Ventriculomegaly

Very Long-Chain Acyl-CoA Dehydrogenase Deficiency (VLCAD)

Very Low Birth Weight <750 gm, <1000 gm

WAGR Syndrome

Walker-Warburg Syndrome

Watson Syndrome

Weaver Syndrome

Weill-Marchesani Syndrome

Werdnig-Hoffman Disease

Wernicke-Korsakoff Syndrome

Williams Syndrome

Wilson Disease

Wolf-Hirschhorn Syndrome

Wolfram Syndrome

Xeroderma Pigmentosum

X-Linked Adrenoleukodystrophy

XXXXX Syndrome

XXYY Syndrome

XYY Syndrome

Zellweger Syndrome