





Established Conditions for the Purposes of 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)	Amyoplasia Congenita	Attention Deficit Hyperactivity Disorder (ADHD)
	Anencephaly	
3-Methylcrotonyl-CoA Carboxylase Deficiency (3MCC)	Angelman Syndrome	Autism Spectrum Disorder (ASD)
	Aniridia-Cerebellar Ataxia Syndrome	Anxiety Disorder
Abusive Head Trauma (previously known as "Shaken Baby Syndrome")	Anopthalmos/Micropthalmos	Baller Gerold Syndrome
		Bannayan Riley Ruvalcaba
Achondrogenesis	Anterior Horn Cell Disease	Bardet-Beidl Syndrome
Acrocallosal Syndrome	Antimongolism Syndrome	
Acrodysotosis	Apert Syndrome	Bartoscas-Papas Syndrome
Acute Disseminated	Antley-Bixler Syndrome	Basal Cell Nevus Syndrome
Encephalomyelitis	APGAR <=3 @20 minutes	Batten Disease
Adrenoleukodystrophy	Aphasia	B-Ketothiolase Deficiency (BKT)
Agenesis of Corpus Callosum		Biotinidase Deficiency (BIOT)
Aicardi Syndrome	Apraxia	Blindness
Aicardi-Goutieres Syndrome	Argininemia (ARG)	Bloch-Sulzberger Syndrome
AIDS	Argininosuccinate Aciduria (ASA)	
	Arhinencephaly	Bobble Head Doll Syndrome
Albinism	Arnold-Chiari Syndrome Type II	Borjeson Syndrome
Alcohol Exposure In Utero	Arthrogryposis	Brain Malformation
Alexander Disease	Ataxia	Brain Tumor
Allan Herndon Syndrome		Brancio-Oto-Renal Syndrome
Alpers Syndrome	Ataxia-Telangiectasia Syndrome	Bulbar Palsy
Alternating Hemiplegia	Attachment Disorders	C Syndrome
Amyelia	Attention Deficit Disorder (ADD)	C Syndrome
Ашуспа		

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Canavan Disease	С
Cancer	C
Carbohydrate Deficient Glycoprotein Syndrome	c c
Carbohydrate Metabolism Disorder	c
Cardio-Facio-Cutaneo Syndrome	c
Carnitine Palmityl Transferase Deficiency Type II (CPTII)	С
Carnitine Uptake Defect/Carnitine Transport Defect (CUD)	c c
Carpenter Syndrome	C
Caudal Regression Syndrome	C
Central Core Disease	C
Cerebellar Agenesis	C
Cerebral Ataxia	C
Cerebral Atrophy	C
Cerebral Degeneration	C
Cerebral Dysgenesis	C
Cerebral/Cortical Dysplasia	C
Cerebral Gigantism	С
Cerebral Palsy	C
Cerebral Venous Thrombosis	C
Cerebrocostomandibular Syndrome	С
Cerebrovascular accident	С
CHARGE Syndrome	С
Chiari Malformation	С
Childhood Disintegrative Disorder	C
Chromosomal Disorders	C
Chromosome 10 Duplication	C
Chromosome 10p+	С
Chromosome 11p-	С

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 2g32 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

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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

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 C_Established Condition List_04-16-2024

Fryns Syndrome

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 C_Established Condition List_04-16-2024 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** C_Established Condition List_04-16-2024

Regulatory Disorders of Sensory	Sjogren-Larsson Syndrome	Timothy Syndrome
Processing	Sly Syndrome	Torticollis
Rett Syndrome		
Rhizomelic Chondrodysplasia Punctata	Small for Gestational Age <=10th percentile	Toxoplasmosis, Congenital
	Smith-Lemli-Opitz Syndrome	Transposition of the Great Vessels
Robert's Syndrome	Smith-Magenis Syndrome	Transverse Myelitis
Rosenberg Chutorian Syndrome		Traumatic Brain Injury
Roussy Levy Syndrome	Social Anxiety Disorder	Trifunctional Protein Deficiency (TFP)
Rubella, Congenital	Sotos Syndrome	Triphosphate Isomerase Deficiency
Rubenstein-Taybi Syndrome	Spastic Paraplegia	Triple X Syndrome
Russel-Silver Syndrome	Sphingolipidoses	Triploidy Syndrome
·	Spielmeyer-Vogt Disease	
Sandhoff Disease	Spina Bifida	Trisomy 12p
Sanfilippo Syndrome	Spinal Cord Injuries	Trisomy 13
Santavuori Disease	Spinal Lipomeningocele	Trisomy 18
Scheie Syndrome	Spinal Muscular Atrophy	Trisomy 3
Schilder's Disease	Spinocerebellar Disorders	Trisomy 8
Schindler Disease		Trisomy 9
Schinzel Giedion Syndrome	Steinert-Myotonic Dystrophy Syndrome	Tuberous Sclerosis
Schizencephaly	Stickler Syndrome	Tyrosinemia Type I and Type II
Schwartz-Jampel Syndrome	Stroke	Urea Cycle Defect
Scott Craniodigital Syndrome	Sturge-Weber Syndrome	Usher Syndrome
Seckel Syndrome	Subacute Sclerosing Panencephalitis	VACTERL Syndrome with
Semilobar Holoprosencephaly	Substance Exposed Infant (SEI)	Hydrocephalus
Shaken Baby Syndrome (now referred	Succinic Semialdehyde	Velocardiofacial Syndrome
to as "Abusive Head Trauma")	Dehydrogenase Deficiency	Ventilator Dependent
Shprintzen-Goldberg Craniosynostosis Syndrome	Sulfatide Lipidosis	Ventriculomegaly
Sialidosis	Sydenham Chorea	Very Long-Chain Acyl-CoA Dehydrogenase Deficiency (VLCAD)
	Syphilis, Congenital	
Sickle Cell Disease/Hemoglobin Disorders (Hb SS, Hb S/B, and Hb S/C)	Syringohydromyelia	Very Low Birth Weight <750 gm, <1000 gm
Simpson Dysmorphia Syndrome	Tay-Sachs Disease	WAGR Syndrome
Sirenomelia Sequence	Tetralogy of Fallot	Walker-Warburg Syndrome
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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