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From:	Birth to Three Early Intervention Program – Administration
Approval Date:	June 7, 2024
Signature :	Erin Rich
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#### **Purpose**

The Birth to Three Early Intervention Program (the Program) maintains an Established Condition List that identifies prevalent physical or mental conditions that would qualify infants and toddlers for Part C services, in accordance with §303.21(a)(2) of the Individuals with Disabilities Education Act. The list is not exhaustive; if an infant or toddler is referred to the Program with a diagnosed physical or mental condition that is not on the Established Condition list, but meets the criteria set forth in §303.21 (2), the infant or toddler may be determined eligible for Part C services.

To appropriately reflect the evolving needs of children and families in Delaware, the established conditions list has been updated and will be implemented effective immediately upon administrative signature and approval.



### **BIRTH MANDATE/Delaware**

	IDEA designations: sensory impairments; severe attachment disorders
F84.0	Autism Spectrum disorder
H47.619	Cortical Blindness, unspecified
H54	Blindness NOS
H54.7	Vision Loss NOS
H90.2	Conductive hearing loss, unspecified
H90.3	Sensorineural Hearing Loss, Bilateral
H90.4	Sensorineural hearing loss, unilateral
H90.41	Sensorineural hearing loss, unilateral, right ear, with unrestricted hearing on the contralateral side
H90.42	Sensorineural hearing loss, unilateral, left ear, with unrestricted hearing on the contralateral side
H90.5	Sensorineural Hearing Loss (Unspec)
H90.71	Mixed conductive and sensorineural hearing loss, unilateral, right ear, with unrestricted hearing on the contralateral side
H90.72	Mixed conductive and sensorineural hearing loss, unilateral, left ear, with unrestricted hearing on the contralateral side
H90.8	Mixed cond & sensorineural hearing loss, unspec
H90.8	Mixed conductive and sensorineural hearing loss, unspecified
H90.A11	Conductive hearing loss, unilateral, right ear with restricted hearing on the contralateral side *
H90.A12	Conductive hearing loss, unilateral, left ear with restricted hearing on the contralateral side *
H90.A2	Sensorineural hearing loss, unilateral, with restricted hearing on the contralateral side
H90.A21	Sensorineural hearing loss, unilateral, right ear, with restricted hearing on the contralateral side
H90.A22	Sensorineural hearing loss, unilateral, left ear, with restricted hearing on the contralateral side
H90.A3	Mixed conductive and sensorineural hearing loss, unilateral with restricted hearing on the contralateral side
H90.A31	Mixed conductive and sensorineural hearing loss, unilateral, right ear with restricted hearing on the contralateral side
H90.A32	Mixed conductive and sensorineural hearing loss, unilateral, left ear with restricted hearing on the contralateral side
H91.2	Sudden idiopathic hearing loss
H91.9	Unspecified hearing loss *
H91.93	Unspecified hearing loss - Bilateral
H93.3X	Disorders of acoustic nerve
H93.3X1	Disorders of right acoustic nerve
H93.3X2	Disorders of left acoustic nerve
H93.3X3	Disorders of bilateral acoustic nerves
H93.3X9	Auditory Neuropathy Spectrum Disorder (ANSD) (Disorder of Acoustic Nerve)
H93.8	Other specified disorders of ear *
H93.8x	Other specified disorders of ear *
H93.8x1	Other specified disorders of right ear *
H93.8x2	Other specified disorders of left ear *

<sup>&</sup>quot;\*" = follow until 12 months post referral – These ICD-10 codes do not always indicate permanent hearing loss.

<sup>&</sup>quot;++" =applies until second birthday



H93.8x3	Other specified disorders of ear, bilateral *
Q16.1	Congenital absence, atresia and stricture of auditory canal (external)

#### MULTIPLE SYSTEM/ GENETIC/ CHROMOSOMAL

_	IDEA designation: chromosomal abnormalities; genetic or congenital disorders; inborn
	errors of metabolism
Q93.5	Angelman Syndrome
Q87.89	Bardet-Biedl Syndrome
Q87.3	Beckwith-Wiedemann Syndrome
Q99.9	Chromosomal Abnormality NOS (Deletion/Dup)
Q99.9	HNRNPU-NDD
Q97.0	Chromosomal Anomalies – Karyotype 47, XXX
E70.0	Classical phenylketonuria
Q90.9	Down Syndrome
Q99.2	Fragile X Syndrome
Q75.4	Franceschetti-Klein (Wildervanck) Syndrome
E75.22	Gaucher Disease (Lipidoses)
Q98.4	Klinefelter's Syndrome
E72.03	Lowe's Syndrome
E71.0	Maple-Syrup-Urine Disease
G71.0	Muscular Dystrophy
Q85.02	Neurofibromatosis type- 2
Q87.19	Noonan Syndrome
Q78.0	Osteogenesis imperfecta
Q87.11	Prader-Willi Syndrome
Q87.0	Saethre–Chotzen syndrome (aka Acrocephalosyndactyly type III)
Q87.2	TAR Syndrome: (Thrombocytopenia with Absent Radius Syndrome)
Q75.4	Treacher Collins Syndrome
Q97.0	Triple X Syndrome (Chromosomal Anom)
Q91.7	Trisomy 13 (Patau's Syndrome) Unspec
Q91.3	Trisomy 18 (Edwards Syndrome) Unspec
Q90.9	Trisomy 21 (Down Syndrome) Unspec
Q85.1	Tuberous sclerosis
Q96.9	Turner-Ullrich Syndrome

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<sup>&</sup>quot;++" =applies until second birthday



# NEUROLOGIC: congenitally or postnatally acquired

Q04.0	Agenesis of Corpus Callosum
G93.1	Anoxic brain damage
Q07.02	Arnold-Chiari syndrome with hydrocephalus
G60.0	Atrophy, Charcot-Marie-tooth Syndrome
163.50	Cerebral Artery Occlusion NOS
G80.8	Cerebral Diplegia / Hemiplegia (congenital)
P10.1	Cerebral Hem. due to birth injury (grade III & IV bleed)
G80.9	Cerebral Palsy, unspecified
G80.0	Cerebral Spastic Quadriplegia
G04.90	Encephalitis (postnatally acquired)
Q07.9	Encephalopathy (congenital)
Q01	Encephalocele
G93.40	Encephalopathies (Degenerative) NOS
D18.02	Hemangioma of intracranial struct.
P91.660	HIE/Body Cooling (Hypoxic Ischemic Enceph.) *
Q04.2	Holoprosencephaly
Q03.8	Hydrocephalus (congenital); Other
Q07.9	Hypomyelination affecting Meningeal bands or folds (congenital)
I61.9	Intracerebral (nontraumatic) Hemorhhage *
G06.0	Intracranial abscess and granuloma
P52.0	Intraventricular Hem./Newborn, only Grade III & IV
E75.4	Jansky-Bielschowsky Amaurotic Idiocy
E75.23	Krabbe disease
G04.81	Leukoencephalitis
P91.2	Leukomalacia, Neonatal Cerebral
Q05.9	Lipomyelomeningocele
Q04.9	Malformation of Brain (Cong/Unspec) *
C71.8	Malig neoplasm of overlapping sites of brain
Q04.5	Megalencephaly
G00.2	Meningitis – Streptococcal*
A02.21	Meningitis due /Salmonella infect.*
G03.9	Meningitis NOS*
Q02	Microcephalus, microcephaly (under 3%)
Q04.3	Microgyria
G70.9	Myoneural disorder
G71.11	Myotonic Dystrophy
Q01.1	Nasofrontal Encephalocele*
E75.4	NCL: Ceroid-Lipofuscinosis, Neuronal



E75.4	NCL: Batten-Mayou Disease
E75.4	NCL: Kuf's Disease (NCL)
H55.01	Nystagmus (congenital)
G91.1	Obstructive Hydrocephalus
H47.039	Optic Nerve Hypoplasia Unspec
Q04.8	Other Specified Congenital Malformation of the Brain *
Q04.3	Polymicrogyria
Q04.6	Porencephalic Cysts (Congenital)
Q04.6	Schizencephaly
Q05.4	Spina Bifida
G12.0	Spinal Muscular Atrophy (Werdnig-Hoffman)
E75.02	Tay Sachs (Nervous System Disorder)
Q93.82	Williams Syndrome

### **NEUROLOGIC - SEIZURES**

P90	Convulsions of Newborn* (only if they are intractable, or with ongoing medication
	treatment)
R56.9	Convulsion NOS* (only if they are intractable, or with ongoing medication treatment)
G40.823	Epil spasms, intractable, w/ status epilepticus
G40.824	Epil spasms, intractable, w/o status epilepticus
G40.919	Epilepsy, unspec, <b>intractable</b> , w/o status epilepticus
G40.911	Epilepsy, unspec, intractable, w/status epilepticus

### **INFECTIONS**

IDEA designation: congenital infections	
P35.1	Cytomegalovirus Infection, Congenital (only if symptomatic)
P35.0	Rubella (Prenatally acquired)
A53.9	Syphilis (Prenatally acquired)
B58.9	Toxoplasmosis (Prenatally acquired)
P39.9	Varicella (Perinatally acquired)
A92.5	Zika Disease only with Microcephaly (not just exposure)

## INFECTIONS: other

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	A48.51	Botulism (Infant)*



## PRENATAL SUBSTANCE EXPOSURE

IDEA de	IDEA designation: disorders secondary to exposure of substances, including fetal alcohol	
	syndrome	
Q86.0	Fetal Alcohol Syn. (dysmorphic, not just alcohol exposure)	
P96.1	NAS - Neonatal Abstinence Syndrome ++	
P96.1	NOWS – Neonatal Opioid Withdrawal Syndrome ++	
P04.1	Phenytoin Exposed (Prenatally)	

#### **CARDIOPULMONARY**

CANDIOI GLINONANI		
Q39.1	Atresia of Esophagus with trachea-esophageal fistula	
P27.1	Bronchopulmonary dysplasia (BPD)* - Only if Oxygen dependent	
l42.8	Cardiomyopathy (Primary)*	
Q25.1	Coarctation of Aorta (pre/postductal)*	
J44.9	Chronic/Obstructive Lung Disease	
Q20.8	Cor Biloculare (Cong. Malf. of Cardiac Chambers and Connections)	
Q20.3	Discordant Ventriculoarterial Connection	
Q20.4	Double inlet ventricle	
Q20.2	Double Outlet left ventricle	
Q20.1	Double outlet right ventricle	
Z92.81	ECMO - Extracorporeal Membrane Oxygenation	
Z94.1	Heart Transplant Status	
l11.0	Hypertensive Heart Disease w/ Heart Failure	
Q23.4	Hypoplastic Left Heart Syndrome	
Q22.6	Hypoplastic Right Heart Syndrome	
Q24.9	Malform of the Heart, Congenital/Unspec*	
I21.3	Myocardial Infarction Unspec (STEMI)	
Z99.81	Oxygen Dependent (Supplemental)	
Q22	Pulmonary Valve Atresia	
Z99.11	Respirator or Ventilator Dependent	
Z93.0	Tracheostomy Status	
Q21.3	Tetralogy of Fallot	
I50.9	Heart Failure *, unspecified	

#### **ENDOCRINE**

E03.1	Hypothyroidism (congenital)*
E23.2	Diabetes Insipidus
E11.9	Diabetes, Diabetic - unspecified
E20.1	Pseudohypoparathyroidism



#### **STRUCTURAL**

Z89.431	Acquired Absence of Right Foot	
Z89.432	Acquired Absence of Left Foot	
Z89.439	Acquired Absence of Unspecified Foot	
Q72.03	Amelia of the Lower Extremities (bilateral)	
S68.412S	Amputation of Left Hand @ Wrist	
S68.411S	Amputation of Right Hand @ Wrist	
Q76.49	Caudel Regression Syndrome	
Q35.9	Cleft Palate* post repair	
Q75.9	Cong Malform of Skull and Face Unspec*	
Q16.4	Congenital Malformation of middle ear NOS	
Q17.9	Congenital Malformations of Ear NOS	
Q31.5	Laryngomalacia	
Q17.2	Microtia	
Q60.2	Renal agenesis, bilateral, unspecified	

Ī	TRAUMA RELATED	
	T74.4	Shaken Infant Syndrome
Ī	S06.2X0D	Traumatic Brain Injury, Diffused w/o LOC

OTHER	
Q74.3	Arthrogryposis
Z94.81	Bone Marrow Transplant*
P07.01	Ex low weight nb, less 500 gr*
P07.02	Ex low weight nb, less 500-749 gr*
P07.03	Ex low weight nb, less 750-999 gr*
P07.26	Less than 28 weeks gestation *
E74.21	Galactosemia*
E74.00	Glycogen storage disorder
E76.02	Hurler-Scheie Syndrome
R20.1	Hypoesthesia of Skin
Z77.011	Lead Exposure (Toxic) If level 5 ug/dl or higher*
P77.9	Necrotizing enterocolitis/NB, unspec* (only if post surgical)
H35.109	Retinopathy of Prematurity, unspec elig., (Grade 3 and above only)
E74.02	Pompe Syndrome
G46.5	PURA syndrome
Q477	Alagille Syndrome *