EarlySteps Eligibility Criteria

ICD-9 to ICD-10 Crosswalk

Purpose: To assist early interventionists in identifying ICD-10 diagnosis codes using the EarlySteps eligibility criteria. The ICD-10 codes will be required for documentation and for billing for Medicaid-paid services for dates of service beginning 10/1/2015. The list below will replace the eligibility criteria list found in Chapter 5 of the Practice Manual. Diagnoses highlighted in blue are additional codes added to the criteria or added to match the "drop down" code list on the eligibility tab used by the SPOEs in EIDS. In addition, the nature of ICD-10 allows for increased specificity in the use of diagnosis coding, therefore the most specific applicable code should always be used. The sequence of the criteria in the list generally follows the sequence in the Practice Manual with some additional categories added to assist in locating codes. If you have questions, please contact your regional coordinator. August, 2015 updates are in red.

Criteria Description	ICD 9	ICD 10	Criteria Description
I. Developmental Delay			
General Category—Specific delays in development	315		
Developmental speech/language disorder	315.3	F80.89	Other developmental disorders of speech and
			language
		F80.9	Developmental disorder of speech and language
			not otherwise specified
Expressive language delay	315.31	F80.1	Expressive language disorder
Mixed receptive and expressive language	315.32	F80.2	Mixed receptive-expressive language disorder
delay		H93.25	Central auditory processing disorder
Speech and language delay due to hearing	315.34	F80.4	Speech and language development delay due to
loss			hearing loss
Developmental Coordination Disorder	315.4	F82	Specific Developmental Disorder: Motor Function
Mixed Developmental Disorder	315.5	F82	Specific Developmental Disorder: Motor Function
Other specified delays in development	315.8	F88	Other disorders of psychological development
II. Use of Informed Clinical Opinion to Determine Eligibility	700.40		
Abnormal sensory-motor response Affective or social disorder/condition	783.42	R62.0	Delayed milestone in childhood
Oral-motor skills dysfunction, including feeding difficulties	783.3	R63.3	Feeding difficulties
III. Established Medical Conditions			
Genetic Disorders			
A. Chromosomal Abnormality Syndromes – General Category	758		
Down syndrome	758.0	Q90.9	Down syndrome, unspecified
Trisomy 13	758.1	Q91.7	Trisomy 13, unspecified (Patau's syndrome)
Trisomy 18	758.2	Q91.3	Trisomy 18, unspecified
General Category Autosomal deletion	758.3_		
syndromes			
Cri-du-chat	758.31	Q93.4	Deletion of short arm of chromosome 5
Velo-cardio-facial syndrome (VCFS)	758.32	Q93.81	Velo-cardio-facial syndrome
Other micro-deletion syndromes: include Miller-Dieker and Smith-Magenis syndromes	758.33	Q93.88	Other microdeletions
DiGeorge Syndrome	279.11	D82.1	Di George's syndrome

Criteria Description	ICD 9	ICD 10	Criteria Description
Fragile X	759.83	Q99.2	Fragile x chromosome
Prader-Willi	759.81	Q87.1	Congenital malformation syndromes
			predominantly associated with short stature
Other conditions due to autosomal	758.5	Q92.8	Other specified trisomies and partial trisomies of
anomalies			autosomes
Other conditions due to chromosomal	758.8_	007.0	1/
anomalies	758.81	Q97.0 Q97.1	Karyotype 47, xxx Female with more than three x chromosomes
Conditions due to sex chromosome		Q97.1 Q97.2	Mosaicism, lines with various numbers of x
anomalies, <u>not</u> including		Q07.2	chromosomes
Klinefelter's Syndrome (XXY)		Q97.8	Other specified sex chromosome abnormalities,
(758.7 [@] Q98.4)or			female phenotype
Turner's syndrome (XO)(758.6 @Q98.9)		Q98.5	Karyotype 47, xyy
		Q98.7	Male with sex chromosome mosaicism
		Q98.8	Other specified sex chromosome abnormalities,
		000.0	male phenotype
Conditions due to enemoly of upon offied	758.9	Q99.8 Q99.9	Other specified chromosome abnormalities Chromosomal abnormality, unspecified
Conditions due to anomaly of unspecified chromosome (includes Williams Syndrome)	758.9	Q99.9	Chromosomal abnormality, unspecified
B. Pre-natal exposures			
Fetal alcohol syndrome	760.71	P04.3	Newborn (suspected to be)affected by maternal
			use of alcohol
		Q86.0	Fetal alcohol syndrome (dysmorphic)
Fetal hydantoin syndrome/Other	760.79	P04.8	Newborn (suspected to be) affected by other
			maternal noxious substances
Narcotics exposure	760.72	P04.49	Newborn (suspected to be) affected by maternal
			use of other drugs of addiction
Hallucinogenic agent exposure	760.73	P04.49	Newborn (suspected to be) affected by maternal
			use of drugs of addiction
Cocaine exposure	760.75	P04.41	Newborn (suspected to be) affected by maternal
-			use of cocaine
Anticonvulsant exposure	760.77	P04.1	Newborn (suspected to be) affected by oth
			maternal medication
Other Noxious influences affecting fetus or	760.79	P04.8	Newborn (suspected to be) affected by other
newborn via placenta or breast milk			maternal noxious substances
Drug Withdrawal Syndrome	779.5	P96.1	Neonatal withdrawal symptoms of maternal use of
	110.0	1 00.1	drugs of addiction
		P96.2	Withdrawal symptoms from therapeutic use of
			drugs of newborn
C. Neurocutaneous Syndromes			
Congenital pigmentary anomalies of the skin	757.33	Q82.1	Xeroderma pigmentosum
	ļ	Q82.2	Mastocytosis
Neurofibromatosis	237.70	Q85.00	Neurofibromatosis, unspecified
Other Neurofibromatosis	237.79	Q85.09	Other neurofibromatosis
Sturge-Weber syndrome	759.6	Q85.8	Other phakomatoses, not elsewhere classified
Tuberous sclerosis	759.5	Q85.1	Tuberous sclerosis
D. Inborn Error of Metabolism	270.0	E70.00	Disordoro of omino poid transport unconstitud
Disorders of amino-acid transport and metabolism	270.0	E72.00 E72.01	Disorders of amino-acid transport, unspecified
III CLADUISIII		E72.01 E72.04	Cystinuria Cystinosis
		E72.04	Other disorders of amino-acid transport
Phenylketonuria (PKU)	270.1	E70.0	Classical phenylketonuria
Other Disturbances of aromatic amino-acid	270.2	E70.21	Tyrosinemia

Criteria Description	ICD 9	ICD 10	Criteria Description
metabolism		E70.29	Other disorders of tyrosine metabolism
		E70.30	Albinism, unspecified
		E70.5	Disorders of tryptophan metabolism
		E70.8	Other disorders of aromatic amino-acid
			metabolism
Maple Sugar Urine Disease	270.3	E71.0	Maple-syrup-urine disease
		E71.120	Methylmalonic acidemia
		E71.19	Other disorders of branched-chain amino-acid
			metabolism
		E71.2	Disorder of branched-chain amino-acid
			metabolism, unspecified
Disturbances of Sulpher-bearing amino acid	270.4	E72.10	Disorders of sulphur-bearing amino-acid
metabolism			metabolism
			Homocystinuria
			Other disorders of sulphur-bearing amino-acid
			metabolism
Disorder of Urea cycle metabolism	270.6	E72.20	Disorder of urea cycle metabolism, unspecified
		E72.22	Arginosuccinic aciduria
		E72.23	Citrullinemia
		E72.29	Other disorders of urea cycle metabolism
Other disturbances of straight-chain amino-	270.7	E72.3	Disorders of lysine and hydroxylysine metabolism
acid metabolism		E72.8	Other specified disorders of amino-acid
			metabolism
Other specified disorders of amino-acid	270.8	E72.03	Lowe's syndrome
metabolism		E72.8	Other specified disorders of amino-acid
			metabolism
Unspecified disorder of amino acid metabolism	270.9	E72.9	Disorder of amino-acid metabolism, unspecified
General Category Disorders of Carbohydrate	271.0	E74.00	Glycogen storage disease, unspecified
Metabolism		E74.01	von Gierke disease
Glycogenosis		E74.04	McArdle disease
		E74.09	Other glycogen storage disease
Galactosemia	271.1	E74.21	Galactosemia
General Category Disorders of Lipid Metabolism	272.0	E78.0	Pure hypercholesterolemia
Lipidoses	272.7		
Fabry's disease		E75.21	Fabry (or Anderson-Fabry) disease
- Gaucher's disease		E75.22	Gaucher disease
- Niemann Pick		E75.249	Niemann-Pick disease, unspecified
- sphingolipidoses		E77.0	Defects in post-translational modification of
			lysosomal enzymes
		E77.1	Defects in glycoprotein degradation
Other disorders of lipid metabolism	272.8	E78.81	Lipoid dermatoarthritis
		E78.89	Other lipoprotein metabolism disorders
		E88.89	Other specified metabolic disorders
Mucopolysaccaridoses	277.5	E76.01	Hurler's syndrome
		E76.03	Scheie's syndrome
		E76.1	Mucopolysaccharidosis, type II-Hunter's syndrome
		E76.219	Morquio mucopolysaccharidoses, unspecified
		E76.22	Sanfilippo mucopolysaccharidoses
		E76.29	Other mucopolysaccharidoses
		E76.3	Mucopolysaccharidosis, unspecified
E. General Category: Cerebral	330		
degenerations of the central nervous			
system—usually manifested in childhood			
Leukodystrophy	330.0	E75.23	Krabbe disease
		E75.25	Metachromatic leukodystrophy
		E75.29	Other sphingolipidosis

Criteria Description	ICD 9	ICD 10	Criteria Description
Cerebral lipidoses such as TaySach's	330.1	E75.02	Tay-Sachs disease
		E75.19	Other gangliosidosis
		E75.4	Neuronal ceroid lipofuscinosis
Cerebral degeneration in generalized lipidoses	330.2	G93.89	Other specified disorders of brain
Cerebral Degenerations of childhood in other diseases	330.3	G93.9	Disorder of brain, unspecified
Other specified degenerations in childhood	330.8	F84.2	Rett's syndrome
		G31.81	Alpers disease
		G31.82	Leigh's disease
Unspecified cerebral degenerations in childhood	330.9	G94	Other disorders of brain in diseases classified elsewhere
F. Prenatal Infections			
TORCH" infections, including:			
Congenital rubella	771.0	P35.0	Congenital rubella syndrome
Congenital cytomegalovirus infection (CMV)	771.1	P35.1	Congenital cytomegalovirus infection
Congenital herpes simplex	771.2	P35.2	Congenital herpesviral [herpes simplex] infection
• · · · · ·		P37.1	Congenital toxoplasmosis
Congenital toxoplasmosis		5050	
		P37.2	Neonatal (disseminated) listeriosis
		P37.8	Other specified congenital infectious and parasitic
C. Other Current areas			diseases
G. Other Syndromes	252.0	F 22.0	A promogoly and nituitory gigantiam
Cerebral gigantism	253.0	E22.0	Acromegaly and pituitary gigantism
General Category: Other and unspecified congenital anomalies	759		
Prader-willi syndrome	759.81	Q87.1	Congenital malformation syndromes
Marfan, and drama	750.00	007.40	predominantly associated with short stature
Marfan syndrome	759.82 759.83	Q87.40 Q99.2	Marfan's syndrome, unspecified
Fragile x syndrome Other specified chromosome abnormalities	759.83	Q99.2 Q99.8	Fragile X chromosome Other specified chromosome abnormalities
Other specified chromosome abnormalities	759.69	E78.71	Barth syndrome
		E78.72	Smith-Lemli-Opitz syndrome
		Q87.1	Cornelia de Lange
		Q87.2	Congenital malformation syndromes
		0.07.12	predominantly involving limbs
		Q87.3	Congenital malformation syndromes involving
			early overgrowth—Beckwith Wiedemann
		Q87.5	Other congenital malformation syndromes with
			other skeletal changes
		Q87.81	Alport syndrome
		Q87.89	Other specified congenital malformation
			syndromes, not elsewhere classified
		Q89.8	Other specified congenital malformations
General Category: Congenital anomaly,	759.9	Q89.9	Congenital anomaly, unspecified
unspecified			
H. Sensory Impairment - Vision			
VisionImpairment can be congenital or	369.00	H54.0	Blindness, both eyes
acquired (369—general category—more			
specific diagnosis obtained from physician)			
Profound impairment, both eyes (369.0-)	200.04	+	
Moderate or severe impairment, better eye,	369.01-		
profound impairment lesser eye Blindness one eye; low vision other eye	260.40		Blindhood, one over low vision other eve
Dimuness one eye, low vision other eye	369.18	H54.10	Blindness, one eye, low vision other eye, unspecified eyes
	200.0	1	
Moderate or severe impairment, both eves	309.2-		
Moderate or severe impairment, both eyes Low vision both eyes not otherwise specified	369.2- 369.20	H54.2	Low vision, both eyes
		H54.2	Low vision, both eyes

Criteria Description	ICD 9	ICD 10	Criteria Description
eye; impairment not further specified	359.24		unspecified eyes
Better eye: moderate vision impairment;	369.25	H54.2	Low vision, both eyes
lesser eye: moderate vision impairment			
Unqualified vision loss, both eyes	369.3	H54.3	Unqualified vision loss, both eyes
Legal blindness, as defined in USA	369.4	H54.8	Legal blindness, as defined in USA
Retrolental fibroplasia or retinopathy of	362.26	H35.159	Retinopathy of prematurity, stage 4, unspecified
prematurity			еуе
ROP Stage 4			
ROP State 5	362.27	H35.169	Retinopathy of prematurity, stage 5, unspecified eye
Bilateral retrolental fibroplasia	362.21	H35.179	Retrolental fibroplasia, unspecified eye
Cortical Blindness	377.75	H47.619	Cortical blindness, unspecified side of brain
I.Sensory Impairment - HearingHearing	389		
impairment (25dB loss or greater) unilateral			
or bilateral General Category			
Conductive hearing loss, unspecified—	389.00	H90.2	Conductive hearing loss, unspecified (includes
includes:			389.00-389.04 and 389.08)
Conductive hearing loss external ear	389.01		
Conductive hearing loss tympanic membrane	389.02		
Conductive hearing loss middle ear	389.03		
Conductive hearing loss inner ear	389.04	1100.44	Openductive backing land, weiletagel vielt approxite
Conductive hearing loss, unilateral	389.05	H90.11	Conductive hearing loss, unilateral, right ear with unrestricted hearing on contralateral side
		H90.12	Conductive hearing loss, unilateral, left ear with unrestricted hearing on contralateral side
Conductive hearing loss, bilateral	389.06	H90.0	Conductive hearing loss, bilateral
Conductive hearing loss of combined types	389.08	H90.2	Conductive hearing loss, unspecified
Sensorineural hearing loss	389.10	H90.5	Unspecified sensorineural hearing loss
Sensory Hearing loss, bilateral	389.11	H903	Sensorineural Hearing loss, bilateral
Neural Hearing loss, bilateral	389.12		
Sensorineural Hearing loss, bilateral	389.18		
Mixed conductive and sensorineural hearing loss	389.20	H90.8	Mixed conductive and sensorineural hearing loss, unspecified
Hearing loss unspecified	389.9	H91.90	Unspecified hearing loss, unspecified ear
Central hearing loss	389.14	H90.5	Unspecified sensorineural hearing loss
J. Orthopedic and Neurological Disorders			
Anoxic brain damage	348.1	G93.1	Anoxic brain damage, not elsewhere classified
Anterior horn cell disease	335.—		
Werdnig-Hoffmann disease	335.0	G12.0	Infantile spinal muscular atrophy, type I [Werdnig- Hoffman]
Spinal muscular atrophy unspecified	335.10	G12.9	Spinal muscular atrophy, unspecified
Kugelberg-welander disease	335.11	G12.1	Other inherited spinal muscular atrophy
Other spinal muscular atrophy	335.19	G12.8	Other spinal muscular atrophies and related syndromes
Amyotrophic lateral sclerosis	335.20	G12.21	Amyotrophic lateral sclerosis
Progressive muscular atrophy	335.21	G12.21	Amyotrophic lateral sclerosis
Progressive bulbar palsy	335.22	G12.22	Progressive bulbar palsy
Pseudobulbar palsy	335.23	G12.8	Other spinal muscular atrophies and related
1		-	syndromes
Primary lateral sclerosis	335.24	G12.29	Other motor neuron disease
Other motor neuron diseases	335.29	G12.29	Other motor neuron disease
Other anterior horn cell diseases	335.8	G12.8	Other spinal muscular atrophies and related syndromes
Anterior horn cell disease unspecified	335.9	G12.9	Spinal muscular atrophy, unspecified
General Category: other specified muscle			,
distorders			
Arthrogryposis	728.3	M62.3	Immobility syndrome (paraplegic)

Criteria Description	ICD 9	ICD 10	Criteria Description
·		M62.89	Other specified disorders of muscle
Arthrogryposis multiplex, congenita	754.89	Q74.3	Arthrogryposis multiplex, congenita
Injury to the Brachial plexus—birth trauma	767.6	P14.0	Erb's paralysis due to birth injury
		P14.1	Klumpke's paralysis due to birth injury
		P14.3	Other brachial plexus birth injuries
Brachial plexus—post perinatal origin	953.4	S14.3XXA	Injury of brachial plexus, initial encounter
Cerebral cysts	348.0	G93.0	Cerebral cysts
Cerebral palsy (all types)- General Category	343		
Congenital diplegia	343.0	G80.1	Spastic diplegic cerebral palsy
Congenital Hemiplegia	343.1	G80.2	Spastic hemiplegic cerebral palsy
Congenital Quadriplegia	343.2	G80.0	Spastic quadriplegic cerebral palsy
Congenital Monoplegia	343.3	G80.8	Other cerebral palsy
Infantile hemiplegia	343.4	G80.2	Spastic hemiplegic cerebral palsy
Other specified infantile cerebral palsy	343.8	G80.8	Other cerebral palsy
Infantile cerebral palsy unspecified	343.9	G80.9	Cerebral palsy, unspecified
Cleft hand	755.58	Q71.60	Lobster-claw hand, unspecified hand
Congenital anomalies of the central	742		•
nervous system – General Category			
Encephalocele	742.0	Q01.9	Encephalocele, unspecified
Microcephaly	742.1	Q02	Microcephaly
Congenital reduction deformities of brain	742.2	Q04.1	Arhinencephaly
5		Q04.2	Holoprosencephaly
		Q04.3	Other reduction deformities of brain
Congenital hydrocephaly	742.3	Q03.0	Malformations of aqueduct of Sylvius
0 7 7 7		Q03.1	Atresia of foramina of Magendie and Luschka
		Q03.8	Other congenital hydrocephalus
Other specified congenital anomalies of	742.4	Q04.5	Megalencephaly
brain		Q04.6	Congenital cerebral cysts
		Q04.8	Other specified congenital malformations of brain
Other specified congenital anomalies of	742.5		
spinal cord—general category			
Diastematomyelia	742.51	Q06.2	Diastematomyelia
Hydromyelia	742.53	Q06.4	Hydromyelia
Other specified congenital anomalies of	742.59	Q06.0	Amyelia
spinal cord		Q06.1	Hypoplasia and dysplasia of spinal cord
		Q06.3	Other congenital cauda equina malformations
		Q06.8	Other specified congenital malformations of spinal
			cord
Other specified congenital anomalies of	742.8	G90.1	Familial dysautonomia [Riley-Day]
nervous system		Q07.8	Other specified congenital malformations of
		_	nervous system
Unspecified congenital anomaly of brain	742.9	Q07.9	Congenital malformation of nervous system,
spinal cord and nervous system			unspecified
Other congenital musculoskeletal	755		
anomalies - General Category			
Reduction of deformities of upper limb	755.20	Q71.899	Other reduction defects of unspecified upper limb
		Q71.90	Unspecified reduction defect of unspecified upper
—		074.00	limb
Transverse deficiency of upper limb	755.21-	Q71.00	Congenital complete absence of unspecified
Longitudinal deficiency of upper limb	755.22	074.10	upper limb
Longitudinal deficiency combined involving	755.23-	Q71.10	Congenital absence of unspecified upper arm and
humerus	755.24	071.05	forearm with hand present.
Longitudinal deficiency , radioulner,	755.25	Q71.20	Congenital absence of both forearm and hand,
complete or partial			unspecified upper limb.
Longitudinal deficiency radial, complete or	755.26	Q71.40	Longitudinal reduction defect of snspecified ratius
partial			

Criteria Description	ICD 9	ICD 10	Criteria Description
Longitudinal deficiency, radial, complete or	755.27	Q71.50	Longitudinal reduction defect of unspecified ulna
partial		L	
Longitudinal deficiency ulnar, carpals or	755.28-	Q71.30	Congenital absence of unspecified hand/finger
metacarpals, phalanges, finger	755.29		
Reduction of deformities of lower limbs	755.30	Q72.899	Other reduction defects of unspecified lower limb
Longitudinal deficiency of lower limb, not	755.32		
classified elsewhere			
Transverse deficiency of lower lime	755.31	Q72.00	Congenital complete absence of unspecified
			lower limb
Longitudinal deficiency combined involving tibia and fibula	755.33	Q72.10	Congenital absence of unspecified thigh and lower leg with foot present
Longitudinal deficiency femoral, complete/incomplete	755.34	Q72.40	Longitudinal reduction defect of unspecified femur
Longitudinal deficiency tibiofibular complete	755.35	Q72.20	Congenital absence of both lower leg and foot,
or partial	755.00	070.50	unspecified lower limb
Longitudinal deficiency, tibia,	755.36	Q72.50	Longitudinal reduction defect of unspecified tibia
complete/partial	755.07	070.00	
Longitudinal deficiency, fibular, complete/partial	755.37	Q72.60	
Longitudinal deficiency, tarsals or	755.38	Q72.30	Congenital absence of unspecified foot and toes
metatarsals complete/partial		Q72.70	Split foot, unspecified lower limb
Longitudinal deficiency, phalanges,	755.39		
complete/partial			
Reduction deformities, unspecified limb	755.4	Q73.0	Congenital absence of unspecified limb(s)
		Q73.1	Phocomelia, unspecified limb(s)
		Q73.8	Other reduction defects of unspecified limb(s)
Congenital cleft hand	755.58	Q71.60	Lobster-claw hand, unspecified hand
Anomalies of skull and face bone	756.0	Q75.0	Craniosynostosis
Premature closure of cranial sutures		Q75.2	Hypertelorism
		Q75.9	Congenital malformation of skull and face bones,
Abaanaa of vortabra, congonital	756.13	Q76.49	unspecified Other congenital malformations of spine, not
Absence of vertebra, congenital	750.15	Q70.49	associated with scoliosis
Chondrodystrophies	756.4	Q77.1	Thanatophoric short stature
		Q77.4	Achondroplasia
		Q77.8	Other osteochondrodysplasia with defects of
			growth of tubular bones and spine
		Q78.4	Enchondromatosis
-			
Osteodystrophies, unspecified	756.50	Q78.9	Osteochondrodysplasia, unspecified
Osteogenesis imperfecta	756.51	Q78.0	Osteogenesis imperfecta
Other symbolic dystunction-general	784.6		
category	704.00	D40.0	Aprovio
Developmental apraxia of speech	784.69	R48.2	Apraxia Other symbolic dysfunctions
Enconholonothy Not Othorwise Operation	240.00	R48.8	Other symbolic dysfunctions
Encephalopathy Not Otherwise Specified	348.30	G93.40	Encephalopathy, unspecified
Fracture of vertebral column with spinal cord	806.00	S12.000A	Unspecified displaced fracture of first cervical
injury (806) General Category—include additional diagnosis from physician		S12.001A	vertebra, initial encounter for closed fracture Unspecified nondisplaced fracture of first cervical
ลินนี้แบบลา นี้เลยาบราร กบกา ยการเปลี่ก		512.001A	vertebra, initial encounter for closed fracture
		S12.100A	Unspecified displaced fracture of second cervical
			vertebra, initial encounter for closed fracture
		S12.101A	Unspecified nondisplaced fracture of second
			cervical vertebra, initial encounter for closed
			fracture
		S12.200A	Unspecified displaced fracture of third cervical
			vertebra, initial encounter for closed fracture
		S12.201A	Unspecified nondisplaced fracture of third cervical
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ICD 9	ICD 10	Criteria Description
		vertebra, initial encounter for closed fracture
	S12.300A	Unspecified displaced fracture of fourth cervical vertebra, initial encounter for closed fracture
	S12.301A	Unspecified nondisplaced fracture of fourth cervical vertebra, initial encounter for closed
	S14.101A	fracture Unspecified injury at C1 level of cervical spinal
	S14.102A	cord, initial encounter Unspecified injury at C2 level of cervical spinal cord, initial encounter
	S14.103A	Unspecified injury at C3 level of cervical spinal cord, initial encounter
	S14.104A	Unspecified injury at C4 level of cervical spinal cord, initial encounter
342		
342.00	G81.00	Flaccid hemiplegia affecting unspecified side
342.01	G81.01	Flaccid hemiplegia affecting right dominant side
	G81.02	Flaccid hemiplegia affecting left dominant side
342.02	G81.03	Flaccid hemiplegia affecting right nondominant side
	G81.04	Flaccid hemiplegia affecting left nondominant side
342.10	G81.10	Spastic hemiplegia affecting unspecified side
342.11	G81.11	Spastic hemiplegia and hemiparesis affecting
		right dominant side
	G81.12	Spastic hemiplegia and hemiparesis affecting left dominant side
		Hemiplegia, unspecified affecting unspecified side
	G81.90	Hemiplegia, unspecified affecting unspecified side
331		
331.3	G91.0	Communicating hydrocephalus
331.4	G91.1	Obstructive hydrocephalus
331.7	G94	Other disorders of brain in diseases classified elsewhere
335.0	G12.0	Infantile spinal muscular atrophy, type I [Werdnig- Hoffman]
331.8	G31.89	Other specified degenerative diseases of the nervous system
345.60	G40.401	Other generalized epilepsy and epileptic syndromes, not intractable, with status epilepticus
	G40.409	Other generalized epilepsy and epileptic syndromes, not intractable, without status
		epilepticus
345.61	G40.411	epilepticus Other generalized epilepsy and epileptic
345.61	G40.411 G40.419	epilepticus
345.61		epilepticus Other generalized epilepsy and epileptic syndromes, intractable, with status epilepticus Other generalized epilepsy and epileptic
345.61 772.13		epilepticus Other generalized epilepsy and epileptic syndromes, intractable, with status epilepticus Other generalized epilepsy and epileptic
	342 342.00 342.01 342.02 342.10 342.11 342.11 342.11 342.11 342.11 342.11 342.11 342.11 342.30 342.90 331 331.3 331.4 331.7 335.0 331.8	S12.300A S12.301A S12.301A S14.101A S14.102A S14.103A S14.103A S14.104A 342 342.00 G81.00 342.01 G81.01 G81.02 342.02 G81.03 G81.04 342.10 G81.10 342.11 G81.11 G81.12 342.80 G81.90 342.90 G81.90 342.90 G81.90 331.3 G91.0 331.4 G91.1 331.7 G94 335.0 G12.0 331.8 G31.89 345.60 G40.401

Criteria Description	ICD 9	ICD 10	Criteria Description
General CategorySpina Bifida/Neural			
Tube Defect	741.00	Q05.4	Unspecified spina bifida with hydrocephalus
		Q07.01	Arnold-Chiari syndrome with spina bifida
		Q07.02	Arnold-Chiari syndrome with hydrocephalus
		Q07.03	Arnold-Chiari syndrome with spina bifida and
			hydrocephalus
Spina Bifida, Cervical region with	741.01	Q05.0	Cervical Spina Bifida with hydrocephalus
Hydrocephalus			
Spina Bifida, dorsal (thoracic) region with	741.2	Q05.1	Thoracic Spina Bidfida with hydrocephalus
hydrcephalus	744.00	0.05.0	
Spina Bifida, lumbar region with	741.03	Q05.2	Lumbar spina bilfida with hydrocephalus
hydrocephalus	741.90	Q05.8	Saaral apina hifida without hydrogophalua
Meningomyelocele		Q05.8 Q05.8	Sacral spina bifida without hydrocephalus
	741.90	Q05.8 Q05.5	Sacral spina bifida without hydrocephalus
Spina Bifida	741.91 741.92		Cervical spina bifida without hydrocephalus
with hydrocephalus	741.92	Q05.6 Q05.7	Thoracic spina bifida without hydrocephalus Lumbar spina bifida without hydrocephalus
General Category: Congenital hereditary	359	Q03.7	
muscular dystrophy	000		
Muscular dystrophies and other myopathies	359.0	G71.2	Congenital myopathies
Hereditary progressive muscular dystrophy	359.1	G71.0	Muscular dystrophy
Myotonic muscular dystrophy	359.21	G71.11	Myotonic muscular dystrophy
Myotonia, congenita	359.22	G71.12	Myotonia congenita
Myontic chondrodystrophy	359.23	G71.13	Myotonic chondrodystrophy
General Category—other paralytic	344		
syndromes			
Quadriplegia and quadriparesisunspecified	344.00	G82.50	Quadriplegia, unspecified
Quadriplegia c1-c4 complete	344.01	G82.51	Quadriplegia, C1-C4 complete
Quadriplegia c1-c4 incomplete	344.02	G82.52	Quadriplegia C1-C4 incomplete
Quadriplegia c5-c7 complete	344.03	G82.53	Quadriplegia, C5-C7 complete
Quadriplegia c5-c7 incomplete	344.04	G82.54	Quadriplegia, C5-C7 incomplete
Other quadriplegia	344.09	G82.50	Quadriplegia, unspecified
Paraplegia	3441	G82.20	Paraplegia, unspecified
Diplegia of upper limbs	344.2	G83.0	Diplegia of upper limbs
Monoplegia of lower limb affecting	344.30	G83.10	Monoplegia of lower limb affecting unspecified
unspecified side			side
Monoplegia of lower limb affecting dominant	344.31	G83.11	Monoplegia of lower limb affecting right dominant
side			side
		G83.12	Monoplegia of lower limb affecting left dominant
Manania si astiawar limb offesting	244.22	002.42	side Manaplacia of lower limb offecting right
Monoplegia of lower limb affecting nondominant side	344.32	G83.13	Monoplegia of lower limb affecting right nondominant side
		G83.14	Monoplegia of lower limb affecting left
		605.14	nondominant side
Monoplegia of upper limb affecting	344.40	G83.20	Monoplegia of upper limb affecting unspecified
unspecified side	344.40	003.20	side
Monoplegia of upper limb affecting dominant	344.41	G83.21	Monoplegia of upper limb affecting right dominant
side	•••••		side
		G83.22	Monoplegia of upper limb affecting left dominant
			side
Monoplegia of upper limb affecting	344.42	G83.23	Monoplegia of upper limb affecting right
nondominant side			nondominant side
		G83.24	Monoplegia of upper limb affecting left
			nondominant side
Unspecified monoplegia	344.5	G83.30	Monoplegia, unspecified affecting unspecified
	244.00	000.4	side
Cauda equina syndrome without neurogenic	344.60	G83.4	Cauda equina syndrome

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Criteria Description	ICD 9	ICD 10	Criteria Description
bladder			
Cauda equina syndrome with neurogenic	344.61	G83.4	Cauda equina syndrome
bladder	244.04	000 5	
Locked-in state Other specified paralytic	344.81 344.89	G83.5 G83.81	Locked-in state Brown-Sequard syndrome
Syndrome	344.09	G83.84	Todd's paralysis (postepileptic)
Syndrome		G83.84 G83.89	Other specified paralytic syndromes
Paralysis unspecified	344.9	G83.9	Paralytic syndrome, unspecified
Paraplegia	344.1	G82.20	Paraplegia, unspecified
Diplegia of upper limbs	344.2	G83.0	Diplegia of upper limbs
Monoplegia of lower limb	344.30	G83.10	Monoplegia of lower limb affecting unspecified
	044.00	000.10	side
Monoplegia of upper limb	344.40	G83.20	Monoplegia of upper limb affecting unspecified
menetre der er offen men			side
Unspecified monoplegia	344.5	G83.30	Monoplegia, unspecified affecting unspecified
			side
General Category— Spinal cord injury	952		
without evidence of spinal bone injury			
Cervical, Dorsal	952.0-		
Range of Codes based on location of injury	952.1	S14.101A-	Unspecified injury at C1 level of cervical spinal
		S34.139A	cord, initial encounter
	0.50.0	0.1.1.00.1	
Unspecified site of spinal cord injury without	952.9	S14.109A	Unspecified injury at unspecified level of cervical
spinal bone injury		004 400 4	spinal cord, initial encounter
		S24.109A	Unspecified injury at unspecified level of thoracic
		S34.109A	spinal cord, initial encounter Unspecified injury at unspecified level of lumbar
		554.109A	spinal cord, initial encounter
		S34.139A	Unspecified injury at unspecified level of sacral
		004.100A	spinal cord, initial encounter
General CategoryOcclusion of cerebral	434		
arteries or stroke			
cerebral thrombosis without cerebral	434.00	166.09	Occlusion and stenosis of unspecified middle
infarction			cerebral artery
		166.19	Occlusion and stenosis of unspecified anterior
			cerebral artery
		166.29	Occlusion and stenosis of unspecified posterior
			cerebral artery
cerebral embolism with cerebral infarction	434.01	163.40	Cerebral infarction due to embolism of
	40.4.4.2		unspecified cerebral artery
Cerebral embolism without cerebral	434.10	166.09	Occlusion and stenosis of unspecified middle
infarction			cerebral artery
		166.19	Occlusion and stenosis of unspecified anterior
		166.20	cerebral artery
		166.29	Occlusion and stenosis of unspecified posterior
		1	cerebral artery
		166 0	
		166.9	Occlusion and stenosis of unspecified cerebral
Cerebral artery occlusion unspecified without	434 00		artery
Cerebral artery occlusion unspecified without	434.90	166.9 166.9	artery Occlusion and stenosis of unspecified cerebral
cerebral infarction		166.9	artery Occlusion and stenosis of unspecified cerebral artery
cerebral infarction General categoryCerebral laceration and	851.00-	I66.9 S06.330A-	arteryOcclusion and stenosis of unspecified cerebral arteryContusion and laceration of cerebrum,
cerebral infarction General categoryCerebral laceration and contusion or traumatic brain injury		166.9	artery Occlusion and stenosis of unspecified cerebral artery Contusion and laceration of cerebrum, unspecified, without loss of consciousness, initial
cerebral infarction General categoryCerebral laceration and	851.00-	I66.9 S06.330A-	arteryOcclusion and stenosis of unspecified cerebral arteryContusion and laceration of cerebrum,

Criteria Description	ICD 9	ICD 10	Criteria Description
Shaken Infant Syndrome	995.55	T74.4XXA	Shaken Infant Syndrome
K. Social Emotional Disorders			
Social Emotional Disorders Childhood Depressive disorders, not elsewhere classified	311	F32.9	Major depressive disorder, single episode, unspecified
Reactive attachment disorder	313.89	F93.8 F94.1 F98.8	Other childhood emotional disorders Reactive attachment disorder of childhood Other specified behavioral and emotional disorders with onset usually occurring in childhood and adolescence
L. Pervasive Developmental Disorders			
Pervasive Developmental Disorders General Category including: Autistic disorder current or active state	299.00	F84.0	Autistic disorder
Autistic disorder current of active state	299.00	F04.0	
Austistice disorder residual state	299.01	F84.0	Autistic disorder
Childhood disintegrative disorder current or active state	299.10	F84.3	Other childhood disintegrative disorder
Childhood disintegrative disorder residual state	299.11	F84.3	Other childhood disintegrative disorder
Other specified pervasive developmental	299.80	F84.5	Asperger's syndrome
disorders current or active state		F84.8	Other pervasive developmental disorders
Other specified pervasive developmental	299.81	F84.5	Asperger's syndrome
disorders residual state		F84.8	Other pervasive developmental disorders
Unspecified pervasive developmental disorder current or active state	299.90	F84.9	Pervasive developmental disorder, unspecified
Unspecified pervasive developmental disorder residual state	299.91	F84.9	Pervasive developmental disorder, unspecified
Asperger syndrome / disorder	299.80	F84.5 F84.8	Asperger's syndrome Other pervasive developmental disorders
M. Medically Related Disorders			
Congenital or infancy-onset hypothyroidism	243	E00.9	Congenital iodine-deficiency syndrome, unspecified
Cleft palate —unspecified	749.00	Q35.9	Cleft palate, unspecified
unilateral, complete	749.01	Q35.9	Cleft palate, unspecified
unilateral, incomplete	749.02	Q35.7	Cleft uvula
	740.40	Q35.9	Cleft palate, unspecified
bilateral, complete	749.13	Q36.0	Cleft lip, bilateral
bilateral, incomplete	749.14	Q36.0 Q37.9	Cleft lip, bilateral
Cleft palate with cleft lip —unspecified unilateral, complete	749.20	Q37.9 Q37.9	Unspecified cleft palate with unilateral cleft lip Unspecified cleft palate with unilateral cleft lip
unilateral, incomplete	749.21	Q37.9 Q37.9	Unspecified cleft palate with unilateral cleft lip
bilateral, complete	749.22	Q37.9 Q37.8	Unspecified cleft palate with bilateral cleft lip
bilateral, incomplete	749.23	Q37.8	Unspecified cleft palate with bilateral cleft lip
Toxic effects of lead and its compounds	984.0	T56.0X1A	Toxic effect of lead and its compounds, accidental
(including fumes) General category	004.0	T56.0X2A	(unintentional), initial encounter Toxic effect of lead and its compounds, intentional self-harm, initial encounter
		T56.0X3A	Toxic effect of lead and its compounds, assault, initial encounter
		T56.0X4A	Toxic effect of lead and its compounds, undetermined, initial encounter
unspecified lead compound effects	984.9	M1A.10X1	Lead-induced chronic gout, unspecified site, with tophus (tophi)
		T56.0X1A	Toxic effect of lead and its compounds, accidental (unintentional), initial encounter

Criteria Description	ICD 9	ICD 10	Criteria Description
		T56.0X2A	Toxic effect of lead and its compounds, intentional
			self-harm, initial encounter
		T56.0X3A	Toxic effect of lead and its compounds, assault,
			initial encounter
		T56.0X4A	Toxic effect of lead and its compounds,
			undetermined, initial encounter
Non-organic failure to thrive	783.41	R62.51	Failure to thrive (child)
Chronic respiratory failure or ventilator	518.83	J96.10	Chronic respiratory failure, unspecified whether
dependence			with hypoxia or hypercapnia
N. Prematurity			
Bronchopulmonary Dysplasia (BPD)	770.7	P27.0	Wilson-Mikity syndrome
		P27.1	Bronchopulmonary dysplasia originating in the
			perinatal period
		P27.8	
Disorders relating to short gestation and low	765.00	P07.00	Extremely low birth weight newborn, unspecified
birth weight General Category—include 5			weight
digit code		P07.10	Other low birth weight newborn, unspecified
			weight
Other preterm infant's birth weight of 1000-	765.10	P07.00	Extremely low birth weight newborn, unspecified
2499 grams —unspecified weight—an			weight
EarlySteps eligibility criterion is <1500 grams		P07.10	Other low birth weight newborn, unspecified
at birth:			weight
less than 500 grams	765.11	P07.01	Extremely low birth weight newborn, less than
			500 grams
500 to 749 grams	765.12	P07.02	Extremely low birth weight newborn, 500-749 g
750 to 999 grams	765.13	P07.03	Extremely low birth weight newborn, 750-999
			grams
1000 to 1249 grams	765.14	P07.14	Other low birth weight newborn, 1000-1249 grams
1250 to 1499 grams	765.15	P07.15	Other low birth weight newborn, 1250-1499 grams
Please refer to weight conversion table at	765.20	P07.20	Extreme immaturity of newborn, unspecified
the end of the chapter.	705.20	F 07.20	weeks of gestation
Weeks of gestation –unspecified gestation-		P07.30	Preterm newborn, unspecified weeks of gestation
-General Category—		107.50	Treterin newborn, unspecified weeks of gestation
EarlySteps eligibility criteria is 32 weeks	765.21	P07.21	Extreme immaturity of newborn, gestational age
gestation or less	705.21	107.21	less than 23 completed weeks
Less than 24 weeks of gestation		P07.22	Extreme immaturity of newborn, gestational age
		1 07.22	23 completed weeks
24 complete weeks of gestation	765.22	P07.23	Extreme immaturity of newborn, gestational age
	1 00.22		24 completed weeks
25-26 weeks of gestation	765.23	P07.24	Extreme immaturity of newborn, gestational age
			25 completed weeks
		P07.25	Extreme immaturity of newborn, gestational age
			26 completed weeks
27-28 weeks of gestation	765.24	P07.26	Extreme immaturity of newborn, gestational age
			27 completed weeks
		P07.31	Preterm newborn, gestational age 28 completed
			weeks
29-30 weeks of gestation	765.25	P07.32	Preterm newborn, gestational age 29 completed
	100.20	1 07.02	weeks
		P07.33	Preterm newborn, gestational age 30 completed
			weeks
31-32 weeks of gestation	765.26	P07.34	Preterm newborn, gestational age 31 completed
	1 00.20		weeks
		P07.35	
		P07.35	Preterm newborn, gestational age 32 completed weeks