



## CMS CLINICAL ELIGIBILITY ATTESTATION

Patient Name: \_\_\_\_\_ DOB: \_\_\_\_\_ Medicaid and/or KidCare ID: \_\_\_\_\_

Parent/Legal Guardian Name: \_\_\_\_\_ Phone number: \_\_\_\_\_

Please note: This form must be completed and attested to by a physician, as defined in Chapter 458 or 459, Florida Statutes. The physician must initial by each applicable ICD10 code. If the code falls within an approved, listed range, the physician must initial and list the specific ICD10 code and descriptor.

Initial all that Apply:

Initials	ICD 10	Descriptor
		<b>B Codes- Viral infections with skin and mucous membrane lesions</b>
	B20	Human immunodeficiency virus (HIV) disease
		<b>C and D Codes- Neoplastic lesions</b>
	C00-D48	All malignancies (except those in remission greater than five years), (list specific ICD10 code and descriptor) _____
		<b>D Codes ≥ 50- Diseases of the blood and blood-forming organs</b>
	D57.00- D57.1	Hb-SS disease, (list specific ICD10 code and descriptor) _____
	D66	Hereditary factor VIII deficiency
	D81.0- D81.4	Severe combined immunodeficiency [SCID], (list specific ICD10 code and descriptor) _____
	D82.1	Digeorge's syndrome
		<b>E Codes- Endocrine, nutritional and metabolic</b>
	E03.1	Congenital Hypothyroidism
	E10.10-E10.9	Type 1 diabetes mellitus, (list specific ICD10 code and descriptor) _____
	E11.00- E11.9	Type 2 diabetes mellitus, (list specific ICD10 code and descriptor) _____
	E25.0	Congenital adrenal hyperplasia
	E70.0	Classical phenylketonuria
	E70.21	Tyrosinemia
	E71.0	Maple Syrup Urine Disease
	E71.110	Isovaleric acidemia
	E71.111	3-methylglutaconic aciduria
	E71.120	Methylmalonic acidemia
	E71.121	Propionic acidemia
	E71.310	LCAD/LCHAD/VLCAD Fatty acid oxidation disorders and ketogenesis disorders
	E71.311	MCAD Fatty acid oxidation disorders and ketogenesis disorders
	E71.312	SCAD Fatty acid oxidation disorders and ketogenesis disorders
	E71.313	Organic Acidurias (glutaric aciduria, Canavan)
	E72.11	Homocystinuria
	E72.21	Argininemia Urea Cycle Disorder

Initials	ICD 10	Descriptor
	E72.22	Arginosuccinic aciduria
	E72.23	Citrulinemia
	E72.4	Ornithine Transcarbamylase Deficiency Urea Cycle Disorder
	E72.51	Non-ketotic Hyperglycinemia
	E74.09	Glycogen storage disease
	E74.21	Galactosemia
	E75.23	Krabbe disease
	E75.240- E75.249	Niemann-Pick disease, (list specific ICD10 code and descriptor) _____
	E75.25	Metachromatic leukodystrophy
	E76.01	Hurler's syndrome
	E76.1	Mucopolysaccharidosis, type II
	E76.22	Sanfilippo mucopolysaccharidoses
	E84.0-84.9	Cystic fibrosis, (list specific ICD10 code and descriptor) _____
		<b>F Codes- Mental, behavioral and neurodevelopmental disorders</b>
	F20.0- F20.9	Schizophrenia (list specific ICD10 code and descriptor) _____
	F25.9	Schizoaffective disorder
	F30.13	Manic episode, severe, without psychotic symptoms
	F30.2	Manic episode, severe with psychotic symptoms
	F31.13	Bipolar disorder, current episode manic without psychotic features, severe
	F31.2	Bipolar disorder, current episode manic severe with psychotic features
	F31.4	Bipolar disorder, current episode depressed, severe, without psychotic features
	F31.5	Bipolar disorder, current episode depressed, severe, with psychotic features
	F31.63	Bipolar disorder, current episode mixed, severe, without psychotic features
	F31.64	Bipolar disorder, current episode mixed, severe, with psychotic features
	F31.81	Bipolar disorder II
	F33.2	Major depressive disorder, recurrent severe without psychotic features
	F33.3	Major depressive disorder, recurrent, severe with psychotic symptoms
		<b>G Codes- Diseases of the nervous system</b>
	G31.81	Alpers disease
	G31.82	Leigh's disease
	G40.001- G40.B19	Epilepsy, (list specific ICD10 code and descriptor) _____
	G71.11	Myotonic muscular dystrophy
	G80.0	Spastic quadriplegia cerebral palsy
	G80.1	Spastic diplegic cerebral palsy
	G80.2	Spastic hemiplegic cerebral palsy
	G82.50- G82.54	Quadriplegia, (list specific ICD10 code and descriptor) _____
	G91.0	Communicating hydrocephalus
	G91.1	Obstructive hydrocephalus
	G93.1	Anoxic brain damage, not elsewhere classified

Initials	ICD 10	Descriptor
		<b>H Codes- H00 to H59 eye and adnexa and H60 to H95 ear and mastoid</b>
	H35.051- H35.059	Retinal neovascularization, (list specific ICD10 code and descriptor) _____
	H35.171- H35.179	Retrolental fibroplasia, (list specific ICD10 code and descriptor) _____
	H54.0	Blindness, binocular
	H90.3	Sensorineural hearing loss, bilateral
	H91.9	Hearing Loss
		<b>I Codes- Diseases of the circulatory system</b>
	I34.0- I34.2	Nonrheumatic mitral (valve), (list specific ICD10 code and descriptor) _____
	I35.0- I35.2	Nonrheumatic aortic (valve), (list specific ICD10 code and descriptor) _____
	I36.1- I36.2	Nonrheumatic tricuspid (valve), (list specific ICD10 code and descriptor) _____
	I50.1-I50.9	All heart failure, (list specific ICD10 code and descriptor) _____
		<b>J Codes- Diseases of the respiratory system</b>
	J45.50- J45.52	Severe persistent asthma, (list specific ICD10 code and descriptor) _____
		<b>K Codes- Diseases of the digestive system</b>
	K72.10	Chronic hepatic failure without coma
	K72.11	Chronic hepatic failure with coma
	K91.2	Short gut syndrome, aka Postsurgical malabsorption
		<b>M Codes- Diseases of the musculoskeletal system and connective tissue</b>
	M08.00- M08.48	Juvenile rheumatoid arthritis, (list specific ICD10 code and descriptor) _____
	M32.0-M32.9	Systemic lupus erythematosus
	M91.11	Juvenile osteochondrosis of head of femur [Legg-Calve-Perthes], right leg
	M91.12	Juvenile osteochondrosis of head of femur [Legg-Calve-Perthes], left leg
		<b>N Codes- Diseases of the genitourinary system</b>
	N18.4	Chronic kidney disease stage IV (severe)
	N18.5	Chronic kidney disease stage V
	N18.6	End stage renal disease
		<b>P Codes- Conditions originating in the perinatal period</b>
	P27.0	Wilson-Mikity syndrome
	P27.1	Bronchopulmonary dysplasia originating in the perinatal period
	P52.21	Intraventricular (nontraumatic) hemorrhage, grade 3, of newborn
	P52.22	Intraventricular (nontraumatic) hemorrhage, grade 4, of newborn
	P78.81	Congenital cirrhosis (of liver)
	P96.0	Congenital renal failure
		<b>Q Codes- Congenital malformations, deformations, and chromosomal abnormalities</b>
	Q00.0	Anencephaly
	Q00.1	Craniorachischisis
	Q00.2	Iniiencephaly
	Q01.0	Frontal encephalocele

Initials	ICD 10	Descriptor
	Q01.1	Nasofrontal encephalocele
	Q01.2	Occipital encephalocele
	Q02	Microcephaly
	Q03.0	Malformations of aqueduct of Sylvius
	Q03.1	Atresia of foramina of Magendie and Luschka
	Q04.2	Holoprosencephaly
	Q04.4	Septo-optic dysplasia of brain
	Q04.5	Megalencephaly
	Q05.0- Q05.3 and Q05.5- Q05.8	Spina bifida, (list specific ICD10 code and descriptor) _____
	Q06.0	Amyelia
	Q06.1	Hypoplasia and dysplasia of spinal cord
	Q07.00- Q07.03	Arnold-Chiari syndrome, (list specific ICD10 code and descriptor) _____
	Q20.0 to Q26.9	Congenital Malformations of the Circulatory System except Peripheral Vascular System Specify Specific ICD-10 Code _____
	Q33.3	Agenesis of lung
	Q33.6	Congenital hypoplasia and dysplasia of lung
	Q35.1	Cleft hard palate
	Q35.3	Cleft soft palate
	Q35.5	Cleft hard palate with cleft soft palate
	Q35.9	Cleft palate, unspecified
	Q36.0	Cleft lip, bilateral
	Q36.1	Cleft lip, median
	Q36.9	Cleft lip, unilateral
	Q37.0	Cleft hard palate with bilateral cleft lip
	Q37.1	Cleft hard palate with unilateral cleft lip
	Q37.2	Cleft soft palate with bilateral cleft lip
	Q37.3	Cleft soft palate with unilateral cleft lip
	Q37.4	Cleft hard and soft palate with bilateral cleft lip
	Q37.5	Cleft hard and soft palate with unilateral cleft lip
	Q37.8	Unspecified cleft palate with bilateral cleft lip
	Q37.9	Unspecified cleft palate with unilateral cleft lip
	Q39.1	Atresia of esophagus with tracheo-esophageal fistula
	Q43.1	Hirschsprung's disease
	Q44.3	Congenital stenosis and stricture of bile ducts
	Q60.1	Renal agenesis, bilateral
	Q75.0	Craniosynostosis
	Q75.1	Craniofacial dysostosis (Crouzan Syndrome)
	Q75.4	Mandibulofacial dysostosis (Treacher Collins)
	Q75.5	Oculomandibular dysostosis (Hallermann-Streif syndrome)
	Q76.1	Klippel-Feil syndrome
	Q78.0	Osteogenesis imperfecta
	Q81.2	Epidermolysis bullosa dystrophica
	Q91.0- Q91.2	Trisomy 18, nonmosaicism, (list specific ICD10 code and descriptor) _____

Initials	ICD 10	Descriptor
	Q91.4-Q91.6	Trisomy 13, (list specific ICD10 code and descriptor) _____
		<b>R Codes- Symptoms, signs and abnormal clinical and laboratory findings, not elsewhere classified</b>
	R40.3	Persistent vegetative state
	R75	Inconclusive laboratory evidence of human immunodeficiency virus (HIV)
		<b>T Codes- Injury, poisoning and certain other consequences of external causes</b>
	T74.1____	Physical abuse, confirmed, (list specific ICD10 code and descriptor) _____
	T74.3____	Psychological abuse, confirmed, (list specific ICD10 code and descriptor) _____
	T74.4XXA- T74.4XXS	Shaken infant syndrome, (list specific ICD10 code and descriptor) _____
		<b>Z Codes- Factors influencing health status</b>
	Z21	Asymptomatic human immunodeficiency virus [HIV] infection status
	Z76.82	Awaiting organ transplant status
	Z94.0	Kidney transplant status
	Z94.1	Heart transplant status
	Z94.2	Lung transplant status
	Z94.3	Heart and lungs transplant status
	Z94.4	Liver transplant status
	Z94.81	Bone marrow transplant status

\*\*\* If you are unable to identify the correct ICD 10 CM diagnosis for your patient that represents a condition that is both chronic and serious with functional limitation, at your option as the patient's physician you may request a Medical Panel Review by providing the following additional information for review after obtaining and including consent from the responsible parent or guardian:

1. Specify the ICD 10 Diagnosis with the Alphabetic Code and up to 5 digits \_\_\_\_\_

Add Additional Diagnoses, if appropriate \_\_\_\_\_

2. Include the most recent specialty consultant summary or hospital discharge summary or a summary of the consideration and description of current condition not to exceed three pages together with the following attestations and Florida licensed physician signature

I certify that the child identified above is actively under my care for the management of the chronic and serious condition(s) identified on this form by my initials.

I certify that my patient is under 21 years of age with a chronic physical, developmental, behavioral, or emotional condition, and also requires health care and related services of a type or amount beyond that which is generally required by children.

I certify my patient has functional limits related to his or her ability to do the things most children of the same age can do.

Patient Name: \_\_\_\_\_ DOB: \_\_\_\_\_

I understand this certification will be used to establish my patient's clinical eligibility for the CMS Managed Care Plan based on the qualifying diagnoses.

Signed: \_\_\_\_\_

Print Name: \_\_\_\_\_

Physician Medicaid #: \_\_\_\_\_

Physician Medical License #: \_\_\_\_\_

Date: \_\_\_\_\_

Office/Practice/Facility Name: \_\_\_\_\_

Office/Practice/Facility Address: \_\_\_\_\_

Office/Practice/Facility Phone Number: \_\_\_\_\_

Office/Practice/Facility Facsimile Number: \_\_\_\_\_

Office/Practice/Facility Contact Person: \_\_\_\_\_

**INSTRUCTIONS: AFTER COMPLETION OF FORM, RETURN TO FLORIDA DEPARTMENT OF HEALTH, OFFICE OF THE CMS MANAGED CARE PLAN VIA EMAIL AT: (CMS.ClinicalEligibilityScreening@flhealth.gov); FACSIMILE (850.488-3813); OR MAIL (Attention: Clinical Eligibility, 4052 Bald Cypress Way, Bin A-06, Tallahassee, Florida, 32399). Parent or guardian acknowledges that emailing is an unsecured method of communication which may result in the unauthorized access of protected health information by third parties.**