

SPECIAL HEALTH SERVICES MEDICAL CONDITIONS Revised: 5-22-2019

- | | |
|--|--|
| <p>ACQUIRED BRAIN INJURY
 ACUTE FLACCID MYELITIS (AFM)
 ADENOID HYPERTROPHY causing SLEEP APNEA
 ALPHA 1-ANTITRYPSIN DEFICIENCY
 AMPUTATION
 AMYOTONIA CONGENITA requiring rehabilitative measures
 ANAL STENOSIS & IMPERFORATE ANUS
 ANEMIAS (excluding minor anemias), including sickle cell
 APLASIA CUTIS CONGENITA, severe, requiring surgery &
 ECTODERMAL DYSPLASIA
 ARNOLD-CHIARI DEFORMITY
 ARTHROGRYPOSIS
 ASTHMA, persistent, requiring controller medications
 ATAXIAS, FAMILIAL DEGENERATIVE DISEASE requiring
 rehabilitative measures
 ATTENTION-DEFICIT/HYPERACTIVITY DISORDER
 (ADD/ADHD)
 AUTO-IMMUNE DISORDERS, chronic, severe, and complex in
 nature</p> <p>BILE DUCT ATRESIA
 BIRTH INJURY (ERB's PALSY, etc.) requiring bracing or
 surgery
 BONE CYST requiring surgery
 BONE TUMORS, benign, requiring surgery, including
 OSTEOCHONDROMAS
 BONY DEFORMITIES requiring bracing, casting or surgery &
 POST-TRAUMATIC DEFORMITY (orthopedic or severe
 soft tissue deformity due to injury; excluding acute fracture
 without an underlying condition)
 BRAIN TUMORS requiring surgery and/or radiation
 BRANCHIOGENIC CLEFT CYST requiring surgery
 BREAST HYPOPLASIA causing considerable psychological
 problems requiring surgery
 BURNS, severe, acute, including residuals</p> <p>CANCER, including CANCER OF EYE
 CATARACTS
 CELIAC DISEASE
 CEREBRAL PALSY, congenital or acquired, requiring
 rehabilitative measures
 CHOANAL ATRESIA
 CHRONIC LUNG/LOWER AIRWAY CONDITIONS, including
 chronic lung disease, chemical pneumonitis, and subglottic
 stenosis
 CLEFT LIP AND/OR PALATE, including SHORT PALATE
 and SUBMUCOUS CLEFT
 COLLAGEN VASCULAR DISORDERS, including but not
 limited to lupus, dermatomyositis, scleroderma, Sjogren's
 syndrome, and rheumatoid arthritis
 CONGENITAL ADRENAL HYPERPLASIA (CAH)
 CORNEAL TRANSPLANTS
 CRANIOSTENOSIS (premature synostosis)
 CYSTIC FIBROSIS
 CYSTIC HYGROMA
 CYSTINOSIS</p> <p>DENTAL DISORDERS, congenital
 DERMATOMYOSITIS
 DIABETES INSIPIDUS</p> | <p>DIABETES MELLITUS, TYPE I and TYPE II
 DIAPHRAGMATIC HERNIA
 DISLOCATION OF HIPS OR OTHER JOINTS</p> <p>EAR DEFORMITY
 EHLERS-DANLOS DISEASE
 ENCEPHALITIS, POLIOMYELITIS OR MENINGITIS,
 residuals of
 ENUCLEATION (removal of eyeball)
 EOSINOPHILIC GASTROENTERITIS
 EPIDERMOLYSIS BULLOSA
 ESOPHAGEAL VARICES
 EYE WOUNDS, penetrating
 EYELID DEFORMITY requiring surgery, congenital</p> <p>FACE DEFORMITY
 FEMORAL CAPITAL EPIPHYSIS, slipped</p> <p>GASTROINTESTINAL TRACT ANOMALIES, congenital
 (including gastroschisis)
 GENITO-URINARY TRACT ANOMALIES, congenital,
 severe and requiring surgery
 GENU RECURVATUM, severe
 GENU VALGUM (Knock-knees), severe
 GENU VARUM (Bowed legs), severe
 GLAUCOMA, congenital
 GROWTH HORMONE DEFICIENCY
 GUILLAIN-BARRE DISEASE, severe, acute, requiring
 tracheotomy and/or ventilation, including residuals</p> <p>HEARING LOSS
 HEART CONDITIONS, congenital or acquired
 HEMANGIOMA, medically significant
 HEMOGLOBINOPATHIES, limited to:
 Sickle cell anemia
 Thalassemia
 HEMOPHILIA including deformities
 HISTIOCYTOSIS X (eosinophilic granuloma)
 HYDROCEPHALUS requiring surgery
 HYPERCHOLESTEROLEMIA, congenital, including familial
 combined hyperlipidemia
 HYPERTHYROIDISM
 HYPOPARATHYROIDISM, congenital or if suspected to last
 longer than two years
 HYPOPHOSPHATEMIC RICKETS
 HYPOPITUITARISM
 HYPOTHALAMIC ADRENAL INSUFFICIENCY
 HYPOTHYROIDISM</p> <p>ICHTHYOSIFORM ERYTHRODERMA, congenital, severe
 IMMUNODEFICIENCY STATES including severe combined
 immunodeficiency (SCID)
 INFLAMMATORY BOWEL DISEASE including Crohn's
 Disease and ulcerative colitis
 INTERSEX DISORDERS, congenital</p> <p>JOINT DEFORMITY, CLUBFEET AND CLUBHANDS,
 severe, requiring bracing, casting, surgery or physical
 therapy</p> |
|--|--|

JUVENILE IDIOPATHIC ARTHRITIS, (Juvenile Rheumatoid Arthritis)

KYPHOSIS, adolescent, requiring bracing or surgery

LARYNGEAL PAPILLOMA

LEUKEMIA

LEUKODYSTROPHY, including adrenoleukodystrophy

MALOCCLUSION, handicapping

MASTOIDITIS, chronic

MEGACOLON requiring surgery

METABOLIC DISORDERS/INBORN ERRORS OF METABOLISM

Amino Acid Disorders, limited to:

- Arginase deficiency/Hyperargininaemia
- Argininemia
- Argininosuccinic acidemia (ASA lyase deficiency)
- Carbamoyl phosphate synthetase deficiency
- Citrullinemia (ASA synthetase deficiency)
- Glutaric acidemia/aciduria
- Glutathione synthetase deficiency (5-oxoprolinuria)
- Homocystinuria (cystathione synthase deficiency)
- Hypermethioninemia
- Hyperornithinemia, hyperammonemia, Homocitrullinemia (HHH syndrome)
- Hyperornithinemia or ornithine oxo-acid aminotransferase deficiency
- Maple syrup urine disease (MSUD)
- N-Acetylglutamate synthetase deficiency
- Nonketotic hyperglycinemia
- Ornithine aminotransferase deficiency
- Ornithine transcarbamylase deficiency (OTC)
- Phenylketonuria (PKU), includes phenylalanine hydroxylase deficiency (PAH) and hyperphenylalaninemia
- Tyrosinemia (I, II, III);

Biotinidase Deficiency

Fatty Acid Oxidation Disorders, limited to:

- 2,4 dienoyl-CoA reductase deficiency
- Long chain acyl-CoA dehydrogenase deficiency (LCADD)
- Long chain 3-OH acyl-CoA dehydrogenase deficiency (LCHAD)
- Carnitine/acylcarnitine translocase deficiency (CACT)
- Carnitine palmitoyltransferase deficiency-type I (CPTI)
- Carnitine palmitoyltransferase deficiency-type II (CPTII)
- Carnitine transport defect (CTD)
- Glutaric acidemia/aciduria
- Medium chain acyl-CoA dehydrogenase deficiency (MCAD)
- Multiple acyl-CoA dehydrogenase deficiency (MADD) or glutaric acidemia-type II (GAII)
- Short chain acyl-CoA dehydrogenase deficiency (SCAD) (ethylmalonic academia)
- Trifunctional protein deficiency (TFP Deficiency)
- Very long chain acyl-CoA dehydrogenase deficiency (VLCAD)

Galactosemia

GLUT 1 Deficiency (glucose 1 transporter deficiency)

Glycogen Storage Disease

Hereditary Fructose Intolerance

Organic Acid Disorders, limited to:

- 2-methylbutyryl-CoA dehydrogenase deficiency
- 3-methylcrotonyl-CoA carboxylase deficiency
- 3-methylglutaconic-CoA hydratase deficiency
- 3-hydroxy-3-methylglutaryl-CoA lyase deficiency
- Glutaric acidemia/aciduria
- Isobutyryl-CoA dehydrogenase deficiency
- Isovaleric acidemia (IVA)
- Methylmalonic acidemia (MMA)
- Propionic Acidemia
- Mitochondrial acetoacetyl-CoA thiolase deficiency (BKT, 3-ketothiolase deficiency)
- Multiple CoA carboxylase deficiency
- Refsum's Disease (Phytanic acid restriction)

MICROCEPHALY

MUCOPOLYSACCHARIDOSIS (MPS) (including variants)

NARCOLEPSY (with or without Cataplexy)

NEPHROSIS & CHRONIC NEPHRITIS

NERVE INJURIES, chronic

NEUROFIBROMATOSIS

NEUROMUSCULAR DISORDERS limited to those covered by MDA including muscular dystrophy

NEVI with malignant potential

OCULAR ALBINISM, congenital

OSTEOCHONDROITIS of various bones

OSTEOGENESIS IMPERFECTA

OSTEOMYELITIS, residuals of

PANTOTHENATE KINASE-ASSOCIATED NEURODEGENERATION (PKAN) (Hallervorden-Spatz Disease, including infusion pump)

PARAPLEGIA, traumatic, and its direct complications

PECTUS CARINATUM/PECTUS EXCAVATUM requiring surgery

PERTHES DISEASE

POLYCYSTIC KIDNEY DISEASE

PRECOCIOUS PUBERTY

PSEUDOHYPOPARATHYROIDISM

PSORIASIS

PTOSIS (drooping eyelids)

PULMONARY LOBAR EMPHYSEMA

RETINAL DETACHMENT in Marfan's syndrome

RETROLENTAL FIBROPLASIA (retinopathy of prematurity)

SCLERODERMA

SCOLIOSIS requiring bracing or surgery

SEIZURE DISORDERS, excluding febrile seizures

SHORT BOWEL SYNDROME

SPINA BIFIDA, MENINGOCELE, MYELOCELE

STRABISMUS through age 10

SUBLUXATED EYE LENS in Marfan's syndrome

SUPERNUMERARY PARTS, severe

SYNDACTYLY

SYNDROMES, limited, requiring ongoing medical treatment; includes septo-optic dysplasia

THROMBOCYTOPENIA, congenital

THROMBOEMBOLISM

THYROGLOSSAL DUCT CYST

T-LYMPHOCYTE IMMUNE DEFICIENCY STATE

TORTICOLLIS (not spasmodic, requiring casting or surgery)

TRACHEAL STENOSIS

TRACHEOESOPHAGEAL FISTULA

TRANSVERSE MYELITIS

TUBERCULOSIS OF BONES AND JOINTS

TUBEROUS SCLEROSIS

UNDESCENDED TESTES

VASCULAR ABNORMALITIES

WEGENER'S GRANULOMATOSIS