

# BABYNET Policy and Procedure Manual

## Appendix 3: Established Risk Conditions

*Children with documentation that any condition on this list has been professionally diagnosed are eligible for BabyNet services based on “established risk”. Families, BabyNet Service Coordinators, and BabyNet Service Providers may link to a guide with a brief description of each condition developed by the Team for Early Childhood Solutions at [http://uscm.med.sc.edu/tecs/babynet\\_covered\\_diagnoses.pdf](http://uscm.med.sc.edu/tecs/babynet_covered_diagnoses.pdf)*

10p13 Deletion	Cataracts w/ Visual Impairment
11q Deletion	Caudal Regression Syndrome
13q Syndrome	Cerebral palsy (CP)/Static Encephalopathy
18q Deletion Syndrome	Charge Association/Syndrome
3Q39	Citrullinemia
49xxxxy syndrome (Multiple x Chromosome Syndrome)	Cleft Hands Bilateral
4p Minus Syndrome	Coffin- Lowry Syndrome
6p Minus Syndrome	Corneal Opacities (Peter’s Anomalie, Sclera cornea)
6q Minus Syndrome	Cornelia de Lange
7q Minus Syndrome	Cortical Vision Impairment
8p Chromosome Deletion	Cri du Chat
Agenesis of the Corpus Callosum	Crouzon Syndrome
Albinism	Cystinosis
Amniotic Band Syndrome	Dandy Walker Malformation
Amyoplasia Congenita Disruptive Sequence	DiGeorge Syndrome
Anencephaly	Down Syndrome (Trisomy 21)
Angelman Syndrome	Duplication Short Arm Chromosome #20
Anophthalmia	Encephalocele
Argininosuccinate lyase deficiency	Fazio-Londe disease
Argininosuccinic Aciduria	Fetal Alcohol Syndrome
Arthrogyriposis	Fragile X
Asphyxia/Hypoxic Ischemic Encephalopathy	Glaucoma w/Visual Impairment
Athetoid Cerebral Palsy	Glutaric Acidemia Type 1
Atresia of the External Auditory Canal	Grade IV Intraventricular Hemorrhage
Auditory Neuropathy	Harlequin Fetus Syndrome
Autism Spectrum Disorder (ASD)	Hearing Loss $\geq$ 20 db
Automatic Eligibility, NOS	Hemiparesis
Bilateral Micromelia	Herpes Encephalitis
Bilateral Optic Nerve Coloboma	Holoprosencephaly
Bilateral Retinal Detachment w/Blindness	Holt Oram Syndrome
Bilateral Visual Acuity $\leq$ 20/70 corrected vision best eye	Hutchinson Gilford Progeria Syndrome
Birthweight $\leq$ 1200 grams or $\leq$ 28 weeks gestational age (until age 2 years)	Hydranencephaly
Brittle Cornea Syndrome (Ehlers-Danlos Syndrome)	Hydrocephaly
Cardio Cranio Cerebello Syndrome	Hypoplastic Left and/or Right Heart Syndrome
Carpenter Syndrome	Incontinentia Pigmenti Syndrome

Infantile Spasms	Pompe Disease
Isochrome 18p Syndrome	Prader-Willi syndrome
Jacobsen's Syndrome	Propionic A acidemia
Joubert Syndrome	R.O.P. stage 4 & 5 Retrolental Fibroplasia
Kabuki syndrome	Retinitis Pigmentosa
Karsch-Neugebauer Syndrome	Retinoblastoma
Klinefelter Syndrome	Rhizomelic Chondrodysplasia Punctata
Krabbe Disease	Ring chromosome 9
Larsen syndrome	Ring chromosome 13
Lebers's Congenital Amaurosis	Rubenstein Taybi Syndrome
Lennox-Gastaut Syndrome	Schizencephaly
Lissencephaly Syndrome	Seckel Syndrome
Lowe Syndrome (oculo-cerebro-renal)	Seizures w/Congenital Brain Malformation
Marshall Smith Syndrome	Septo-Optic Dysplasia
Melnick-Frazier	Severe Attachment Disorder (ASD)
Methylmalonic Acidemia	Shaken Baby Syndrome/Abusive Head Trauma
Microdactyly	Smith-Magenis Syndrome
Microtia	Spastic Diplegia
Midas Syndrome	Spastic Hemiplegia
Miller-Dieker Syndrome	Spastic Quadriplegia
Mobius sequence or Mobius Syndrome	Spina Bifida
MPS (Mucopolysaccharidosis)	Spinal Cord Injury
MSUD (Maple Syrup Urine Disease)	Spinal Muscular Atrophy
Myelodysplasia	Stickler Syndrome
Myotonic Dystrophy	Syringohydromyelia
Myotubular Myopathy	Tar syndrome
Non-Ketotic Hyperglycemia	Tay - Sachs disease
Neural Tube Defects	Tetrasomy 12p
Ohtahara Syndrome	Treacher Collins Syndrome
Opitz Syndrome	Trisomy 1
Optic Nerve Atrophy	Trisomy 5p
Ornithine-Carbamyl-Transferase Deficiency	Trisomy 10
Osteogenesis Imperfecta	Trisomy 13
Osteopetrosis	Trisomy 18
P1 Deletion Chromosome	Trisomy 4
Pachygyria	Trisomy 8 Mosaicism Syndrome
Pallister-Killian syndrome	Trisomy 9
Pathologic Head Growth	Tuberous Sclerosis
Perinatal Asphyxia, severe	Turner's syndrome
Pervasive Developmental Disorder (ASD)	Usher Syndrome
Phocomelia	Vater Syndrome, with Limb Anomalies
Pierre Robin Syndrome/Sequence	Velo-Cardo-Facial Syndrome
PKU	Waardenberg Syndrome

Werdnig-Hoffman  
William's Syndrome  
Wolf-Hirschhorn Syndrome  
Zellweger Spectrum Syndrome