## **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

10p13 Deletion 11q Deletion 13q Syndrome

18q Deletion Syndrome

3Q39

49xxxxy syndrome (Multiple x Chromosome Syndrome)

4p Minus Syndrome 6p Minus Syndrome 6q Minus Syndrome 7q Minus Syndrome 8p Chromosome Deletion

Agenesis of the Corpus Callosum

Albinism

Amniotic Band Syndrome

Amyoplasia Congenita Disruptive Sequence

Anencephaly

Angelman Syndrome Anophthalmia

Argininosuccinate lyase deficiency

Argininosuccinic Aciduria

Arthrogryposis

Asphyxia/Hypoxic Ischemic Encephalopathy

Athetoid Cerebral Palsy

Atresia of the External Auditory Canal

**Auditory Neuropathy** 

Autism Spectrum Disorder (ASD)

Automatic Eligibility, NOS Bilateral Micromelia

Bilateral Optic Nerve Coloboma

Bilateral Retinal Detachment w/Blindness

Bilateral Visual Acuity ≤ 20/70 corrected vision best eye

Birthweight  $\leq$  1200 grams or  $\leq$  28 weeks gestational age

(until age 2 years)

Brittle Cornea Syndrome (Ehlers-Danlos Syndrome)

Cardio Cranio Cerebello Syndrome

Carpenter Syndrome

Cataracts w/ Visual Impairment
Caudal Regression Syndrome

Cerebral palsy (CP)/Static Encephalopathy

Charge Association/Syndrome

Citrullinemia

Cleft Hands Bilateral Coffin- Lowry Syndrome

Corneal Opacities (Peter's Anomalie, Sclera

cornea)

Cornelia de Lange

Cortical Vision Impairment

Cri du Chat

Crouzon Syndrome

Cystinosis

Dandy Walker Malformation

DiGeorge Syndrome

Down Syndrome (Trisomy 21)

Duplication Short Arm Chromosome #20

Encephalocele

Fazio-Londe disease Fetal Alcohol Syndrome

Fragile X

Glaucoma w/Visual Impairment

Glutaric Acidemia Type 1

Grade IV Intraventricular Hemorrhage

Harlequin Fetus Syndrome

Hearing Loss ≥ 20 db

Hemiparesis

Herpes Encephalitis Holoprosencephaly Holt Oram Syndrome

Hutchinson Gilford Progeria Syndrome

Hydranencephaly Hydrocephaly

Hypoplastic Left and/or Right Heart 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

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 SyndromeSchizencephalyLissencephaly SyndromeSeckel Syndrome

Lowe Syndrome (oculo-cerebro-renal) Seizures w/Congenital Brain Malformation

Marshal Smith Syndrome Septo-Optic Dysplasia

Melnick-Frazier Severe Attachment Disorder (ASD)

Methylmalonic Acidemia Shaken Baby Syndrome/Abusive Head Trauma

Microdactyly Smith–Magenis Syndrome

MicrotiaSpastic DiplegiaMidas SyndromeSpastic HemiplegiaMiller-Dieker SyndromeSpastic 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