BIRTH DEFECTS AND GENETIC DISEASES BRANCH 6-DIGIT CODE

For Reportable Congenital Anomalies

Based on the 1979 British Pediatric Association (BPA) Classification of Diseases and the World Health Organization's 1979 International Classification of Diseases, 9th Revision, Clinical Modification (ICD-9-CM)

Code modifications developed by Division of Birth Defects and Developmental Disabilities, National Center on Birth Defects and Developmental Disabilities, Centers for Disease Control and Prevention, Public Health Service, U.S. Department of Health and Human Services, Atlanta, Georgia 30333

Doc. No. 6digit88
Version 08/07
Replaces Versions 06/04, 05/07, and 06/07

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Explanation of 6-Digit Code

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6th Digit Code - Master

.000 Blank
.001 Left Only
.002 Right Only
.003 Unilateral Unspecified
.004 Bilateral
.005
.006
.007
.008 Possible, Probable, Borderline, or Rule Out;
Defects only diagnosed prenatally should be coded with the last digit 8 when the prenatal diagnosis is not definitive.
.009 Not Otherwise Specified (NOS)
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Notes:

An asterisk (*) beside a disease code indicates that the code was created by CDC.

A pound symbol (#) beside a disease code indicates that the condition or defect is listed on the MACDP Exclusion List.

A check (T) beside a disease code indicates that an addition/revision was made since the last printing of the Procedure Manual. Use of the code should be according to the exclusion list criteria.

The abbreviations NEC and NOS used in this code are defined as not elsewhere classified and as not otherwise specified, respectively.

CONGENITAL ANOMALIES

Anencephalus and Similar Anomalies

740.0 Anencephalus

740.000 Absence of brain 740.010 Acrania 740.020 Anencephaly 740.030 Hemianencephaly, hemicephaly 740.080 Other

740.1 Craniorachischisis

740.100 Craniorachischisis

740.2 Iniencephaly

740.200 Closed iniencephaly740.210 Open iniencephaly740.290 Unspecified iniencephaly

741 Spina Bifida

741.0 Spina Bifida with Hydrocephalus

741.000 Spina bifida aperta, any site, with hydrocephalus 741.010 Spina bifida cystica, any site, with hydrocephalus and Arnold-Chiari malformation Arnold-Chiari malformation, NOS 741.020 Spina bifida cystica, any site, with stenosed aqueduct of Sylvius 741.030 Spina bifida cystica, cervical, with unspecified hydrocephalus Spina bifida cystica, cervical, with hydrocephalus but without mention of Arnold-Chiari malformation or aqueduct stenosis 741.040 Spina bifida cystica, thoracic, with unspecified hydrocephalus, no mention of Arnold-Chiari 741.050 Spina bifida cystica, lumbar, with unspecified hydrocephalus, no mention of Arnold-Chiari 741.060 Spina bifida cystica, sacral, with unspecified hydrocephalus, no mention of Arnold-Chiari 741.070 Spina bifida of any site with hydrocephalus of late onset

- 741.080 Other spina bifida, meningocele of specified site with hydrocephalus 741.085 Spina bifida, meningocele, cervicothoracic, with hydrocephalus 741.086 Spina bifida, meningocele thoracolumbar, with hydrocephalus 741.087 Spina bifida, meningocele, lumbosacral with hydrocephalus 741.090 Spina bifida of any unspecified type
- with hydrocephalus

741.9 Spina bifida without mention of hydrocephalus

- 741.900 Spina bifida (aperta), without hydrocephalus 741.910 Spina bifida (cystica), cervical, without hydrocephalus 741.920 Spina bifida (cystica), thoracic, without hydrocephalus 741.930 Spina bifida (cystica), lumbar, without hydrocephalus 741.940 Spina bifida (cystica), sacral, without hydrocephalus 741.980 Spina bifida, other specified site, without hydrocephalus
- Includes: cervicothoracic, thoracolumbar, lumbosacral
- 741.985 Lipomyelomeningocele
- 741.990 Spina bifida, site unspecified, without hydrocephalus (myelocoele, myelomeningocele, meningomyelocele)

742 Other Congenital Anomalies of Nervous System

742.0 Encephalocele

- 742.000 Occipital encephalocele 742.080 Other encephalocele of specified site (includes midline defects) 742.085 Frontal encephalocele 742.086 Parietal encephalocele 742.090 Unspecified encephalocele
- 742.1 Microcephalus

742.100 Microcephalus

742.2 Reduction deformities of brain

- 742.200 Anomalies of cerebrum
- 742.210 Anomalies of corpus callosum
- 742.220 Anomalies of hypothalamus
- 742.230 Anomalies of cerebellum
- 742.240 Agyria and lissencephaly
- 742.250 Microgyria, polymicrogyria
- 742.260 Holoprosencephaly
- 742.270 Arrhinencephaly
- 742.280 Other specified reduction defect of brain
- 742.290 Unspecified reduction defect of brain

742.3 Congenital hydrocephalus

Excludes: hydrocephalus with any condition in 741.9 (use 741.0)

- 742.300 Anomalies of aqueduct of Sylvius
- 742.310 Atresia of foramina of Magendie and Luschka

| # | 742.380 | Dandy-Walker syndrome Hydranencephaly Other specified hydrocephaly Includes: communicating hydrocephaly Hydrocephalus secondary to intraventricular hemorrhage (IVH) or CNS bleed |
|-------|--|---|
| | 742.390 | Unspecified hydrocephaly, NOS |
| 742.4 | Other spe | ecified anomalies of brain |
| | 742.400 | Enlarged brain and/or head megalencephaly macrocephaly |
| | 742.410 | Porencephaly Includes: porencephalic cysts |
| | 742.420 | Cerebral cysts |
| | | Other specified anomalies of brain |
| | | Includes: cortical atrophy cranial nerve defects |
| | 742.485 | Ventricular cysts |
| | | Excludes: arachnoid cysts |
| | 742.486 | Small brain |
| 742.5 | 5 Other specified anomalies of spinal cord | |
| | 742.500 | Amyelia |
| | 742.510 | Hypoplasia and dysplasia of spinal cord |
| | | atelomyelia |
| | | myelodysplasia |
| | 742.520 | Diastematomyelia |
| | | Other cauda equina anomalies |
| | 742.540 | Hydromyelia |
| | | Hydrorachis |
| | 742.580 | Other specified anomalies of spinal cord and membranes Includes: congenital tethered cord |
| 742.8 | | ecified anomalies of nervous system |
| | Excludes | congenital oculofacial paralysis: |
| | | Moebius syndrome (use 352.600) |
| | 742.800 | Jaw-winking syndrome Marcus Gunn syndrome |
| | 742.810 | Familial dysautonomia |
| | 742.010 | Riley-Day syndrome |
| | 742.880 | Other specified anomalies of nervous system |
| 742.9 | Unspecifi | ied anomalies of brain, spinal cord and nervous systems |
| | 742.900 | Brain, unspecified anomalies |
| | 742.910 | Spinal cord, unspecified anomalies |
| | 742.990 | Nervous system, unspecified anomalies |

743 Congenital Anomalies of Eye

743.000 Anophthalmos agenesis of eye cryptophthalmos
743.100 Microphthalmos, small eyes aplasia of eye hypoplasia of eye dysplasia of eye rudimentary eye

743.2 Buphthalmos

743.200 Buphthalmos
congenital glaucoma
hydrophthalmos
743.210 Enlarged eye, NOS
743.220 Enlarged cornea
keratoglobus
congenital megalocornea

743.3 Congenital cataract and lens anomalies

743.300 Absence of lens
congenital aphakia

743.310 Spherical lens
Spherophakia

743.320 Cataract, NOS

743.325 Cataract, anterior polar

743.326 Cataract, other specified

743.330 Displaced lens

743.340 Coloboma of lens

743.380 Other specified lens anomalies

743.390 Unspecified lens anomalies

743.4 Coloboma and other anomalies of anterior segments

743.400 Corneal opacity 743.410 Other corneal anomalies Excludes: megalocornea (use 743.220) 743.420 Absence of iris aniridia 743.430 Coloboma of iris 743.440 Other anomalies of iris polycoria ectopic pupil Peter's anomaly Excludes: brushfield spots (use 743.800) 743.450 Blue sclera If <36 weeks gestation, code only if another reportable defect is present. Always code if ≥36 weeks gestation. 743.480 Other specified colobomas and anomalies of anterior segments Rieger's anomaly 743.490 Unspecified colobomas and anomalies of anterior eye segments

743.5 Congenital anomalies of posterior segment

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743.500 Specified anomalies of vitreous humour
743.510 Specified anomalies of retina
congenital retinal aneurysm
Excludes: Stickler syndrome (use 759.860)

743.520 Specified anomalies of optic disc
hypoplastic optic nerve
coloboma of the optic disc

743.530 Specified anomalies of choroid
743.535 Coloboma of choroid
743.536 Other specified anomalies of posterior segment of eye
743.590 Unspecified anomalies of posterior segment of eye
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743.6 Congenital anomalies of eyelids, lacrimal system, and orbit

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743.600 Blepharoptosis
                congenital ptosis
       743.610 Ectropion
       743.620 Entropion
    # 743.630 Other anomalies of eyelids
                absence of eyelashes
                long eyelashes
                weakness of eyelids
Т
                fused eyelids (exclude if <25 weeks gestation unless another
                reportable defect is present)
       743.635 Blepharophimosis
                small or narrow palpebral fissures
       743.636 Coloboma of the eyelids
       743.640 Absence or agenesis of lacrimal apparatus
                absence of punctum lacrimale
      743.650 Stenosis or stricture of lacrimal duct
       743.660 Other anomalies of lacrimal apparatus (e.g., cyst) 743.670 Anomalies of orbit
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743.8 Other specified anomalies of eye

* 743.810 Epibulbar dermoid cyst

743.9 Unspecified anomalies of eye

743.900 Unspecified anomalies of eye congenital: of eye (any part) anomaly, NOS deformity, NOS

744 Congenital Anomalies of Ear, Face, and Neck

744.0 Anomalies of ear causing impairment of hearing

- 744.000 Absence or stricture of auditory canal
- 744.010 Absence of auricle (pinna)

absence of ear, NOS

744.020 Anomaly of middle ear

fusion of ossicles

744.030 Anomaly of inner ear

Includes: congenital anomaly of membranous

labyrinth organ of Corti

744.090 Unspecified anomalies of ear with hearing impairment

Includes: congenital deafness, NOS

744.1 Accessory auricle

744.100 Accessory auricle

Polyotia

744.110 Preauricular appendage, tag, or lobule

(in front of ear canal)

744.120 Other appendage, tag, or lobule include papillomas, ear tags

744.2 Other specified anomalies of ear

- 744.200 Macrotia (enlarged pinna)
- 744.210 Microtia (hypoplastic pinna and absence or stricture of external auditory meatus)
- 744.220 Bat ear
- Т # 744.230 Other misshapen ear

pointed ear

elfin

pixie-like

lop ear

cauliflower ear

cleft in ear

malformed ear

absent or decreased cartilage

- 744.240 Misplaced ears
- 744.245 Low set ears 744.246 Posteriorly rotated ears
 - 744.250 Absence or anomaly of eustachian tube
- 744.280 Other specified anomalies of ear (see also 744.230)

Excludes: Darwin's tubercle

744.3 Unspecified anomalies of ear

744.300 Unspecified anomalies of ear Congenital: ear (any part) anomaly, deformity, NOS

744.4 Branchial cleft, cyst, or fistula; preauricular sinus

744.400 Branchial cleft, sinus, fistula cyst, or pit # 744.410 Preauricular sinus, cyst, or pit

744.480 Other branchial cleft anomalies Includes: dermal sinus of head

744.500 Webbing of neck

Includes: pterygium colli,

redundant neck skin folds

744.8 Other unspecified anomalies of face and neck

- 744.800 Macrostomia (large mouth) 744.810 Microstomia (small mouth) # 744.820 Macrocheilia (large lips)
 # 744.830 Microcheilia (small lips)
 744.880 Other specified anomalies of face/neck

744.9 Unspecified anomalies of face and neck

- # 744.900 Congenital anomaly of neck, NOS Includes: short neck
 - 744.910 Congenital anomaly of face, NOS Abnormal facies

745 Bulbus Cordis Anomalies and Anomalies of Cardiac Septal Closure

745.0 Common truncus (see 747.200 for pseudotruncus)

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745.000 Persistent truncus arteriosus
absent septum between aorta and pulmonary
artery
745.010 Aortic septal defect
Includes: aortopulmonary window
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Excludes: <u>atrial</u> septal defect (use 745.590)

745.1 Transposition of great vessels

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745.100 Transposition of great vessels, complete (no VSD)
       745.110 Transposition of great vessels, incomplete (w/ VSD)
               Taussig-Bing syndrome
       745.120 Corrected transposition of great vessels,
               L-transposition, ventri in version
               Excludes: dextrocardia (use 746.800)
      745.130 Double outlet right ventricle (DORV) with normally
               related great vessels
      745.140 Double outlet right ventricle (DORV) with transposed
               great vessels
      745.150 Double outlet right ventricle (DORV), relationship of great
N
               vessels not specified
      745.180 Other specified transposition of great vessels,
N
               no mention of double outlet right ventricle (DORV)
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745.190 Unspecified transposition of great vessels

745.2 Tetralogy of Fallot

```
745.200 Fallot's tetralogy
745.210 Fallot's pentalogy
Fallot's tetralogy plus ASD
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745.3 Single ventricle

745.300 Single ventricle
Common ventricle
Cor triloculare biatriatum

745.4 Ventricular septal defect

```
745.400 Roger's disease
               Note: This is an oudated term and the code is no longer
                      used. If this diagnostic term is encountered in
                      the medical record, code it as a ventricular septal
                      defect.
      745.410 Eisenmenger's syndrome
      745.420 Gerbode defect
Т
      745.480 Other specified ventricular septal defect
               Includes: cystalline
                           sub-cystalline
                           subarterial
                           conoventricular
      745.485
              Perimembranous VSD
               Includes: membranous VSD
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745.486
        Muscular VSD
         Includes: mid-muscular and apical VSDs
745.487
         Inlet VSD
         Includes: common atrioventricular (AV) canal type VSD
         Note: Code common atrioventricular (AV) canal as
                745.630
                Code common atrioventricular (AV) canal with
                muscular VSD as 745.620
         Ventricular septal defect, NOS
745.490
         Excludes: common atrioventricular canal type (use
         745.620)
745.498
        Probable VSD
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745.5 Ostium secundum type atrial septal defect

| N | # 745.500 | Nonclosure of foramen ovale, NOS Patent foramen ovale (PFO) 1) Always code if ≥36 weeks of gestation at birth and defect last noted at ≥6 weeks of age. 2) If ≥36 weeks gestation at birth and defect last noted <6 weeks of age, code only if another reportable heart defect is present. 3) Never code if <36 weeks gestation at birth regardless of presence of other defects. |
|---|--------------------|--|
| S | 745.510 | Ostium (septum) secundum defect Note: If the defect size by echo is <= 4mm, assume it is a PFO and follow the coding instructions for 745.500, even if the record says secundum ASD. |
| N | 745.520 | Lutembacher's syndrome Note: This is an outdated term and the code is no longer used. If this diagnostic term is encountered in the medical record, code the individual components, not the syndrome. |
| S | 745.570 | |
| S | 745.580 745.590 | Other specified atrial septal defect ASD (atrial or auricular septal defect), NOS Excludes: PFO vs. ASD (see 745.570). Note: If the defect size by echo is <= 4mm, assume it is a PFO and follow the coding instructions for 745.500, even if the record says ASD. |

745.6 Endocardial cushion defects

745.600 Ostium primum defects
745.610 Single common atrium, cor triloculare biventriculare
N 745.620 Common atrioventricular canal with ventricular septal defect (VSD)
Includes: Common AV canal with muscular VSD

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Excludes: Inlet VSD or common AV canal type VSD (code as
                    745.487
           745.630 Common atrioventricular canal
           745.680 Other specified cushion defect
           745.690 Endocardial cushion defect, NOS
    745.7 Cor biloculare
           745.700 Cor biloculare
    745.8 Other specified defects of septal closure
           745.800 Other specified defects of septal closure
    745.9 Unspecified defect of septal closure
           745.900 Unspecified defect of septal closure
    746
          Other Congenital Anomalies of Heart
    746.0 Anomalies of pulmonary valve
           746.000 Atresia, hypoplasia of pulmonary valve
                    Note: Code pulmonary artery atresia as 747.300
                           Code pulmonary artery hypoplasia as 747.380
                           Code "pulmonic" or "pulmonary" atresia or hypoplasia, NOS (no mention of valve or
                           artery) as 746.995
    Ν
           746.010 Stenosis of pulmonary valve
                    Excludes:
                                  pulmonary infundibular
                                  stenosis (use 746.830)
                           Code pulmonary artery stenosis as 747.320
                           Code "pulmonic" or "pulmonary" stenosis, NOS (no
                           mention of valve or artery) as 746.995
    N
        # 746.020
                   Pulmonary valve insufficiency or regurgitation,
                    congenital
                    Never code cases designated as 'mild', minimal', 'trivial',
                    or 'physiologic'.
                    Code cases designated as 'moderate' or 'severe' and those
                    where the degree is not specified (NOS) only if another
                    reportable heart defect is present.
           746.080 Other specified anomalies of pulmonary valve
                                  pulmonary infundibular
                    Excludes:
                                  stenosis (use 746.830)
           746.090 Unspecified anomaly of pulmonary valve
    746.1 Tricuspid atresia and stenosis
           746.100 Tricuspid atresia only
                    Excludes: tricuspid stenosis and hypoplasia
        # 746.105 Tricuspid valve insufficiency or regurgitation,
                    congenital
                    Never code cases designated as 'mild', minimal', 'trivial',
                    or 'physiologic'.
                    Code cases designated as 'moderate' or 'severe' and those
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N 746.106 Tricuspid stenosis or hypoplasia

746.2 Ebstein's anomaly

746.200 Ebstein's anomaly

746.3 Congenital stenosis of aortic valve

746.300 Congenital stenosis of aortic valve

Excludes: supravalvular aortic stenosis (747.220)

746.4 Congenital insufficiency of aortic valve

N # 746.400 Aortic valve insufficiency or regurgitation, congenital Excludes: bicuspid aortic valve.

Never code cases designated as 'mild', minimal', 'trivial',

or 'physiologic'.

Code cases designated as 'moderate' or 'severe' and those where the degree is not specified (NOS) only if another

reportable heart defect is present.

N 746.470 Bicuspid aortic valve

* 746.480 Other specified anomalies of the aortic valves

Includes: aortic valve atresia

Excludes: <u>supra</u>valvular aortic stenosis (747.220)

* 746.490 Unspecified anomalies of the aortic valves

746.5 Congenital mitral stenosis

746.500 Congenital mitral stenosis

746.505 Absence, atresia, or hypoplasia of mitral valve

746.6 Mitral valve insufficiency or regurgitation, congenital

N # 746.600 Mitral valve insufficiency or regurgitation, congenital Never code cases designated as 'mild', minimal', 'trivial', or 'physiologic'. Code cases designated as 'moderate' or 'severe' and those where the degree is not specified (NOS) only if another

746.7 Hypoplastic left heart syndrome

746.700 Hypoplastic left heart syndrome
Atresia, or marked hypoplasia of the
ascending aorta and defective development
of left ventricle (with mitral valve atresia)

reportable heart defect is present.

746.8 Other specified anomalies of the heart

746.800 Dextrocardia without situs inversus (situs solitus)
Dextrocardia with no mention of situs inversus
Excludes: dextrocardia with situs inversus use 759.300)

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746.810 Levocardia
               Note: This condition has been moved to the never code
                      list.
      746.820 Cor triatriatum
      746.830 Pulmonary infundibular (subvalvular) stenosis
      746.840 Trilogy of Fallot
      746.850 Anomalies of pericardium
    # 746.860 Anomalies of myocardium
Ν
               cardiomegaly, congenital, NOS
               cardiomyopathy, congenital
               cardiomyopathy, hypertrophic
               Note: Do not code cardiomyopathy of any type in a
               newborn of a diabetic mother (either gestational or pre-
               existing diabetes).
              Congenital heart block
      746.870
      746.880 Other specified anomalies of heart
               Includes:
                             ectopia (ectopic) cordis (mesocardia),
                             conduction defects, NOS
      746.881 Hypoplastic left ventricle
               Excludes:
                            hypoplastic left heart syndrome (746.700)
      746.882 Hypoplastic right heart (ventricle)
               Uhl's disease
      746.883 Hypoplastic ventricle, NOS
      746.885 Anomalies of coronary artery or sinus
      746.886 Ventricular hypertrophy (right or left)
Ν
               Note: Do not code ventricular hypertrophy of any type in
               a newborn of a diabetic mother (either gestational or
               pre-existing diabetes).
      746.887 Other defects of the atria
                             congenital Wolfe-Parkinson-White
               Excludes:
                             (use 426.705)
                             rhythm anomalies (use 426.-, 427.-)
746.9 Unspecified anomalies of heart
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746.900 Unspecified anomalies of heart valves
      746.910 Anomalous bands of heart
      746.920 Acyanotic congenital heart disease, NOS
      746.930 Cyanotic congenital heart disease, NOS
               Blue baby
      746.990 Unspecified anomaly of heart:
                Includes: congenital heart disease (CHD)
      746.995
               "Pulmonic" or "pulmonary" atresia, stenosis, or
N
               hypoplasia, NOS (no mention of valve or artery)
                      Code pulmonary valve atresia or hypoplasia as
               Note:
                       746.000
                      Code pulmonary valve stenosis as 746.010
                      Code pulmonary artery atresia as 747.300
                       Code pulmonary artery stenosis as 747.320
                       Code pulmonary artery hypoplasia as 747.380
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747 Other Congenital Anomalies of Circulatory System

2) If ≥36 weeks gestation at birth and defect last noted <6 weeks of age, code only if the PDA was treated (e.g. by ligation or indomethicin) or if another reportable heart defect is present.

3) Never code if <36 weeks gestation at birth or if treated with prostaglandins regardless of gestational age. (See PDA Tree Appendix) Probable PDA

747.1 Coarctation of aorta

747.008

- 747.100 Preductal (proximal) coarctation of aorta 747.110 Postductal (distal) coarctation of aorta 747.190 Unspecified coarctation of aorta
- 747.2 Other anomalies of aorta
- 747.200 Atresia of aorta absence of aorta pseudotruncus arteriosus 747.210 Hypoplasia of aorta tubular hypoplasia of aorta 747.215 Interrupted aortic arch, Type A N 747.216 Interrupted aortic arch, Type B N 747.217 Interrupted aortic arch, Type C 747.220 Supra-aortic stenosis (supravalvular) Excludes: aortic stenosis, congenital (see 746.300) 747.230 Persistent right aortic arch 747.240 Aneurysm of sinus of Valsalva 747.250 Vascular ring (aorta) double aortic arch Includes: vascular ring compression of trachea 747.260 Overriding aorta dextroposition of aorta 747.270 Congenital aneurysm of aorta congenital dilatation of aorta 747.280 Other specified anomalies of aorta 747.285 Interrupted aortic arch, NOS, type not specified 747.290 Unspecified anomalies of aorta

747.3 Anomalies of pulmonary artery

Pulmonary artery atresia, absence or agenesis 747.300 Note: Code pulmonary valve atresia as 746.000 Code "pulmonic" or "pulmonary" atresia, NOS (no mention of valve or artery) as 746.995 747.310 Pulmonary artery atresia with septal defect 747.320 Pulmonary artery stenosis N Includes: Stenosis of the main pulmonary artery or of the right or left main branches Code pulmonary valve stenosis as 746.010 Code "pulmonic" or "pulmonary" stenosis, NOS (no mention of valve or artery) as 746.995 747.325 Ν Peripheral pulmonary artery stenosis Includes: Stenosis of a pulmonary artery peripheral to

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the main right or left main branches
                          Peripheral pulmonic stenosis (PPS), NOS,
                          documented by echocardiogram
                Excludes: Peripheral pulmonic stenosis (PPS) murmur only
                (not documented by echocardiogram) Note: 1) Always code if \geq 36 weeks of gestation at birth and
                defect last noted at \geq 6 weeks of age.
                2)If ≥36 weeks gestation at birth and defect last noted
                <6 weeks of age, code only if another reportable heart
                defect is present.
                3) Never code if <36 weeks gestation at birth.
                (See PPS Tree Appendix)
      747.330 Aneurysm of pulmonary artery
                dilatation of pulmonary artery
      747.340 Pulmonary arteriovenous malformation or aneurysm
      747.380 Other specified anomaly of pulmonary artery
                             pulmonary artery hypoplasia
                Includes:
                Note: Code pulmonary valve hypoplasia as 746.000
                       Code "pulmonic" or "pulmonary" hypoplasia, NOS
                       (no mention of valve or artery) as 746.995
      747.390 Unspecified anomaly of pulmonary artery
747.4 Anomalies of great veins
      747.400 Stenosis of vena cava (inferior or superior)
      747.410 Persistent left superior vena cava
      747.420 (TAPVR) Total anomalous pulmonary venous return
      747.430 Partial anomalous pulmonary venous return
      747.440 Anomalous portal vein termination
      747.450 Portal vein - hepatic artery fistula
      747.480 Other specified anomalies of great veins
      747.490 Unspecified anomalies of great veins
747.5 Absence or hypoplasia of umbilical artery
    # 747.500 Single umbilical artery
747.6 Other anomalies of peripheral vascular system
      747.600 Stenosis of renal artery
      747.610 Other anomalies of renal artery
      747.620 Arteriovenous malformation (peripheral)
                Excludes: pulmonary (747.340) cerebral (747.800)
                          retinal (743.510)
      747.630 Congenital phlebectasia
                congenital varix
      747.640 Other anomalies of peripheral arteries
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aberrant subclavian artery

Includes: primary pulmonary artery hypertension ONLY if

it is present in an infant at ≥7 days of age
747.690 Unspecified anomalies of peripheral vascular system

Budd-Chiari - occlusion of hepatic vein (use

S = Rev. 8/07 R = Rev. 6/07 N = Rev. 5/07 T = Rev. 6/04

N # 453.000)
747.680 Other anomalies of peripheral vascular system

747.650 Other anomalies of peripheral veins

Includes:

Excludes:

747.8 Other specified anomalies of circulatory system

747.9 Unspecified anomalies of circulatory system

747.900 Unspecified anomalies of circulatory system

748 Congenital Anomalies of Respiratory System

748.0 Choanal atresia

748.000 Choanal atresia atresia of nares, anterior or posterior congenital stenosis

748.1 Other anomalies of nose

```
748.100 Agenesis or underdevelopment of nose
748.110 Accessory nose
748.120 Fissured, notched, or cleft nose
748.130 Sinus wall anomalies
748.140 Perforated nasal septum

# 748.180 Other specified anomalies of nose
flat bridge of nose
wide nasal bridge
small nose and nostril
absent nasal septum

748.185 Tubular nose, single nostril, proboscis
748.190 Unspecified anomalies of nose
Excludes: congenital deviation of the nasal
septum (use 754.020)
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748.2 Web of larynx

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748.205 Web of larynx-glottic
748.206 Web of larynx-subglottic
748.209 Web of larynx-NOS
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748.3 Other anomalies of larynx, trachea, and bronchus

748.300 Anomalies of larynx and supporting cartilage
748.310 Congenital subglottic stenosis - Never code if chart states the condition was acquired or secondary to endotracheal (ET) intubation or ventilation
748.330 Other anomalies of trachea

Excludes: vascular ring compression of the trachea (use 747.250)

748.340 Stenosis of bronchus
748.350 Other anomalies of bronchus
748.360 Congenital laryngeal stridor, NOS
748.380 Other specified anomalies of larynx and bronchus
748.385 Cleft larynx, laryngotracheoesophageal cleft
748.390 Unspecified anomalies of larynx, trachea, and bronchus

748.4 Congenital cystic lung

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748.400 Single cyst, lung or lung cyst
748.410 Multiple cysts, lung
Polycystic lung
748.420 Honeycomb lung
748.480 Other specified congenital cystic lung
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748.5 Agenesis or aplasia of lung

748.500 Agenesis or aplasia of lung

T 748.510 Hypoplasia of lung; Pulmonary hypoplasia

Exclude if isolated defect in infants <36 weeks gestation.

748.520 Sequestration of lung

748.580 Other specified dysplasia of lung

Fusion of lobes of lung

* 748.590 Unspecified dysplasia of lung

748.6 Other anomalies of lung

748.600 Ectopic tissues in lung
748.610 Bronchiectasis
748.620 Accessory lobe of lung
748.625 Bilobar right lung or right lung with left lung bronchial pattern
748.690 Other and unspecified anomalies of lung

748.8 Other specified anomalies of respiratory system

748.800 Anomaly of pleura
748.810 Congenital cyst of mediastinum
748.880 Other specified respiratory system anomalies
Includes: congenital lobar emphysema
lymphangiectasia of lungs

748.9 Unspecified anomalies of respiratory system

748.900 Unspecified anomalies of respiratory system
Absence of respiratory organ, NOS
Anomaly of respiratory system, NOS

749 Cleft Palate and Cleft Lip

749.0 Cleft palate alone

(If description of condition includes Pierre Robin sequence, use additional code, 524.080)

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749.000 Cleft hard palate, unilateral
749.010 Cleft hard palate, bilateral
749.020 Cleft hard palate, central
749.030 Cleft hard palate, NOS
749.040 Cleft soft palate, alone unilateral
749.050 Cleft soft palate, alone bilateral
749.060 Cleft soft palate, alone central
749.070 Cleft soft palate, alone, NOS
749.080 Cleft uvula
749.090 Cleft palate, NOS
palatoschisis
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749.1 Cleft lip alone

749.2 Cleft lip with cleft palate

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749.200 Cleft lip, unilateral, with any cleft palate 749.210 Cleft lip, bilateral, with any cleft palate 749.220 Cleft lip, central, with any cleft palate 749.290 Cleft lip, NOS, with any cleft palate
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750 Other Congenital Anomalies of Upper Alimentary Tract

750.000 Tonque tie Ankyloglossia

750.1 Other anomalies of tongue

Excludes:protruding tongue (never a defect)

750.100 Aglossia Absence of tongue 750.110 Hypoglossia (small tongue) Microglossia 750.120 Macroglossia (large tongue) 750.130 Dislocation or displacement of tongue Glossoptosis 750.140 Cleft tongue or split tongue

750.180 Other specified anomalies of tongue 750.190 Unspecified anomalies of tongue

750.2 Other specified anomalies of mouth and pharynx

750.200 Pharyngeal pouch 750.210 Other pharyngeal anomalies 750.230 Other anomalies of salivary glands or ducts 750.240 High arched palate 750.250 Other anomalies of palate 750.260 Lip fistulae or pits 750.270 Other lip anomalies Includes: notched lip, prominent philtrum, long philtrum cleft lip (see 749) Excludes: 750.280 Other specified anomalies of mouth and pharynx Excludes: receding jaw (see 524.0)

750.3 Tracheoesophageal (T-E) fistula, esophageal atresia and stenosis

750.300 Esophageal atresia without mention of T-E fistula 750.310 Esophageal atresia with mention of T-E fistula 750.320 Tracheoesophageal fistula without mention of esophageal atresia 750.325 Tracheoesophageal fistula - "H" type 750.330 Bronchoesophageal fistula with or without mention of esophageal atresia 750.340 Stenosis or stricture of esophagus 750.350 Esophageal web 750.380 Other tracheoesophageal anomalies

large and small mouth (see 744.8)

750.4 Other specified anomalies of esophagus

750.400 Congenital dilatation of esophagus giant esophagus 750.410 Displacement of esophagus 750.420 Diverticulum of esophagus esophageal pouch 750.430 Duplication of esophagus 750.480 Other specified anomalies of esophagus

750.5 Congenital hypertrophic pyloric stenosis

- # 750.500 Pylorospasm
 750.510 Congenital hypertrophic pyloric stenosis
 750.580 Other congenital pyloric obstruction
- 750.6 Congenital hiatus hernia
 - 750.600 Congenital hiatus hernia
 Cardia displacement through esophageal hiatus
 Partial thoracic stomach
 Excludes: congenital diaphragmatic hernia (756.610)

750.7 Other specified anomalies of stomach

- 750.700 Microgastria
 750.710 Megalogastria
 750.720 Cardiospasm
 achalasia of cardia, congenital
 750.730 Displacement or transposition of stomach
 750.740 Diverticulum of stomach
 750.750 Duplication of stomach
 750.780 Other specified anomalies of stomach
- 750.8 Other specified anomalies of upper alimentary tract

750.800 Other specified anomalies of upper alimentary tract

750.9 Unspecified anomalies of upper alimentary tract

750.900 Unspecified anomalies of mouth and pharynx 750.910 Unspecified anomalies of esophagus 750.920 Unspecified anomalies of stomach 750.990 Unspecified anomalies of upper alimentary tract

751 Other Congenital Anomalies of Digestive System

751.0 Meckel's diverticulum

- 751.000 Persistent omphalomesenteric duct persistent vitelline duct
- # 751.010 Meckel's diverticulum

751.1 Atresia and stenosis of small intestine

- 751.100 Stenosis, atresia or absence of duodenum
- 751.110 Stenosis, atresia or absence of jejunum
- 751.120 Stenosis, atresia or absence of ileum
- 751.190 Stenosis, atresia or absence of small intestine
- 751.195 Stenosis, atresia or absence of small intestine with fistula

751.2 Atresia and stenosis of large intestine, rectum and anal canal

- 751.200 Stenosis, atresia or absence of large intestine Stenosis, atresia or absence of appendix
- 751.210 Stenosis, atresia or absence of rectum with fistula
- 751.220 Stenosis, atresia or absence of rectum without mention of fistula
- 751.230 Stenosis, atresia or absence of anus with fistula Includes: imperforate anus with fistula
- 751.240 Stenosis, atresia or absence of anus without mention of fistula Includes: imperforate anus without fistula

751.3 Hirschsprung's disease and other congenital functional disorders of the colon

- 751.300 Total intestinal aganglionosis
- 751.310 Long-segment Hirschsprung's disease; aganglionosis beyond the rectum
- 751.320 Short-segment Hirschsprung's disease; aganglionosis involving no more than the anal sphincter and the rectum
- 751.330 Hirschsprung's disease, NOS
- 751.340 Congenital megacolon congenital macrocolon, not aganglionic

751.4 Anomalies of intestinal fixation

- 751.400 Malrotation of cecum and/or colon
- 751.410 Anomalies of mesentery
- 751.420 Congenital adhesions or bands of omentum and peritoneum; Ladd's bands
- 751.490 Other specified and unspecified malrotation
- 751.495 Malrotation of small intestine alone

751.5 Other anomalies of intestine

- 751.500 Duplication of anus, appendix, cecum, or intestine enterogenous cyst
- 751.510 Transposition of appendix, colon, or intestine
- 751.520 Microcolon

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751.530 Ectopic (displaced) anus
       751.540 Congenital anal fistula
       751.550 Persistent cloaca
R
       751.555 Exstrophy of cloaca
                Excludes exstrophy of urinary bladder not associated with
                imperforate anus (use 753.500)
    * 751.560
               Duodenal web
    # 751.580 Other specified anomalies of intestine
                Includes: rectal fissures
       751.590 Unspecified anomalies of intestine
751.6 Anomalies of gallbladder, bile ducts, and liver
       751.600 Absence or agenesis of liver, total or partial
       751.610 Cystic or fibrocystic disease of liver
    # 751.620 Other anomalies of liver
                hepatomegaly
                hepatosplenomegaly (also use code 759.020)
                              Budd-Chiari (use 453.000)
                Excludes:
       751.630 Agenesis or hypoplasia of gallbladder
       751.640 Other anomalies of gallbladder
                duplication of gallbladder
       751.650 Agenesis or atresia of hepatic or bile ducts
                Includes:
                             biliary atresia
                Excludes:
                              congenital or neonatal hepatitis
                              (use 774.480 or 774.490)
      751.660 Choledochal cysts
751.670 Other anomalies of hepatic or bile ducts
751.680 Anomalies of biliary tract, NEC
751.7 Anomalies of pancreas
       Excludes: fibrocystic disease of pancreas (277.000)
                diabetes mellitus,
                congenital
                neonatal
       751.700 Absence, agenesis or hypoplasia of pancreas
       751.710 Accessory pancreas
       751.720 Annular pancreas
       751.730 Ectopic pancreas
       751.740 Pancreatic cyst
       751.780 Other specified anomalies of pancreas
       751.790 Unspecified anomalies of pancreas
751.8 Other specified anomalies of digestive system
       751.800 Absence of alimentary tract, NOS
                (complete or partial)
       751.810 Duplication of alimentary tract
       751.820 Ectopic digestive organs, NOS
       751.880 Other specified anomalies of digestive system
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751.9 Unspecified anomalies of digestive system

751.900 Unspecified anomalies of digestive system congenital of digestive system, NOS anomaly, NOS deformity, NOS obstruction, NOS

752 Congenital Anomalies of Genital Organs

Excludes: congenital hydrocele (778.600) testicular feminization syndrome (257.800) syndromes associated with anomalies in number and form of chromosomes (758)

752.0 Anomalies of ovaries

- 752.000 Absence or agenesis of ovaries
- 752.010 Streak ovary
- 752.010 Stream Ovary
- 752.080 Other specified anomalies of ovaries 752.085 Multiple ovarian cysts 752.090 Unspecified anomalies of ovaries

752.1 Anomalies of fallopian tubes and broad ligaments

- 752.100 Absence of fallopian tube or broad ligament
- 752.110 Cyst of mesenteric remnant epoophoron cyst

cyst of Gartner's duct

- 752.120 Fimbrial cyst
 - parovarian cyst
- 752.190 Other and unspecified anomalies of fallopian tube and broad ligaments

752.2 Doubling of uterus

752.200 Doubling of uterus doubling of uterus (any degree) or associated with doubling of cervix and vagina

752.3 Other anomalies of uterus

- 752.300 Absence or agenesis of uterus
- 752.310 Displaced uterus
- 752.320 Fistulae involving uterus with digestive or urinary tract

uterointestinal fistula Includes: uterovesical fistula

- 752.380 Other anomalies of uterus
 - bicornuate uterus unicornis uterus
- 752.390 Unspecified anomalies of uterus

752.4 Anomalies of cervix, vagina, and external female genitalia

- 752.400 Absence, atresia or agenesis of cervix
- 752.410 Absence or atresia of vagina, complete or partial
- 752.420 Congenital rectovaginal fistula
- 752.430 Imperforate hymen
- # 752.440 Absence or other anomaly of vulva

fusion of vulva

hypoplastic labia majora - Always code if $\geq 36 weeks$ gestation. If <36 weeks gestation, code only if another reportable defect is present.

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# 752.450 Absence or other anomaly of clitoris
Includes: clitoromegaly
enlarged clitoris
clitoral hypertrophy
prominent clitoris

# 752.460 Embryonal cyst of vagina
752.470 Other cyst of vagina, vulva, or canal of Nuck
0ther specified anomalies of cervix, vagina, or external female
genitalia
Includes: vaginal tags
hymenal tags
```

752.490 Unspecified anomalies of cervix, vagina, or external female genitalia

752.5 Undescended testicle

1)If < 36 weeks gestation, code only if there is a medical/surgical intervention for this problem; 2)If \(\geq 36 \) weeks gestation and defect last noted at <1 year of age, code only if there was a medical/surgical intervention for this problem or if another reportable defect is present 3)Always code if \(\geq 36 \) weeks gestation and defect first noted at \(\geq 1 \) of age.

752.514 Undescended testicle, bilateral

752.520 Undescended testicle, NOS (Cryptorchidism) 752.530 Ectopic testis, unilateral and bilateral

752.6 Hypospadias and epispadias

752.600 Hypospadias (alone), NOS
752.605 1°, glandular,coronal
752.606 2°, penile
752.607 3°, perineal, scrotal
752.610 Epispadias
752.620 Congenital chordee (with hypospadias), NOS
752.621 Congenital chordee alone (chordee w/o hypospadias)
752.625 Cong. chordee with 1°, coronal hypospadias
752.626 Cong. chordee with 2°, penile hypospadias
752.627 Cong. chordee with 3°, perineal, scrotal hypospadias

752.7 Indeterminate sex and pseudohermaphroditism

Excludes:pseudohermaphroditism:

female, with adrenocortical disorder (see 255.200) male, with gonadal disorder with specified chromosomal anomaly (see 758)

752.700 True hermaphroditism ovotestis

752.710 Pseudohermaphroditism, male
752.720 Pseudohermaphroditism, female
pure gonadal dysgenesis
Excludes: gonadal agenesis (758.690)
752.730 Pseudohermaphrodite, NOS
752.790 Indeterminate sex, NOS
ambiguous genitalia

752.8 Other specified anomalies of male genital organs

752.800 Absence of testis monorchidism, NOS # 752.810 Aplasia or hypoplasia of testis and scrotum 752.820 Other anomalies of testis and scrotum polyorchidism bifid scrotum Excludes: torsion of the testes or spermatic cord (use #608.200) 752.830 Atresia of vas deferens 752.840 Other anomalies of vas deferens and prostate 752.850 Absence or aplasia of penis 752.860 Other anomalies of penis absent or hooded foreskin redundant foreskin (never a defect) 752.865 Small penis, hypoplastic penis, or micropenis 752.870 Cysts of embryonic remnants cyst: hydatid of Morgagni Wolffian duct appendix testis 752.880 Other specified anomalies of genital organs microgenitalia macrogenitalia

752.9 Unspecified anomalies of genital organs

752.900 Unspecified anomalies of genital organs Congenital: of genital organ, NEC anomaly, NOS or deformity, NOS

753 Congenital Anomalies of Urinary System

753.0 Renal agenesis and dysgenesis

- 753.000 Bilateral absence, agenesis, dysplasia, or hypoplasia of kidneys Potter's syndrome 753.009 Renal agenesis, NOS
- 753.010 Unilateral absence, agenesis, dysplasia or hypoplasia of kidneys

753.1 Cystic kidney disease

- 753.100 Renal cyst (single)
- 753.110 Polycystic kidneys, infantile type
- 753.120 Polycystic kidneys, adult type
- 753.130 Polycystic kidneys, NOS
- 753.140 Medullary cystic disease, juvenile type
- 753.150 Medullary cystic disease, adult type
- Medullary sponge kidney 753.160 Multicystic renal dysplasia
- Multicystic kidney
- 753.180 Other specified cystic disease Includes: cystic kidneys, NOS

753.2 Obstructive defects of renal pelvis and ureter

- 753.200 Congenital hydronephrosis
- 753.210 Atresia, stricture, or stenosis of ureter

ureteropelvic junction obstruction/stenosis Includes: ureterovesical junction obstruction/stenosis hypoplastic ureter

- 753.220 Megaloureter, NOS
 - hydroureter Includes:
- 753.290 Other and unspecified obstructive defects of renal pelvis and ureter

753.3 Other specified anomalies of kidney

- 753.300 Accessory kidney
- 753.310 Double or triple kidney and pelvis

pyelon duplex or triplex

- 753.320 Lobulated, fused, or horseshoe kidney
- 753.330 Ectopic kidney
- 753.340 Enlarged, hyperplastic or giant kidney
- 753.350 Congenital renal calculi
- 753.380 Other specified anomalies of kidney

753.4 Other specified anomalies of ureter

- 753.400 Absence of ureter
- 753.410 Accessory ureter

double ureter, duplex collecting system

- 753.420 Ectopic ureter
- 753.480 Other specified anomalies of ureter Includes: ureterocele

753.5 Exstrophy of urinary bladder

753.500 Exstrophy of urinary bladder ectopia vesicae extroversion of bladder

753.6 Atresia and stenosis of urethra and bladder neck

- 753.600 Congenital posterior urethral valves or posterior urethral obstruction
- 753.610 Other atresia, or stenosis of bladder neck
- 753.620 Obstruction, atresia or stenosis of anterior urethra
- 753.630 Obstruction, atresia or stenosis of urinary meatus Includes: meatal stenosis
- 753.690 Other and unspecified atresia and stenosis of urethra and bladder neck

753.7 Anomalies of urachus

- T # 753.700 Patent urachus
 - 753.710 Cyst of urachus
 - 753.790 Other and unspecified anomaly of urachus

753.8 Other specified anomalies of bladder and urethra

- 753.800 Absence of bladder or urethra
- 753.810 Ectopic bladder
- 753.820 Congenital diverticulum or hernia of bladder
- 753.830 Congenital prolapse of bladder (mucosa)
- 753.840 Double urethra or urinary meatus
- 753.850 Ectopic urethra or urethral orifice
- 753.860 Congenital digestive-urinary tract fistulae rectovesical fistula
- 753.870 Urethral fistula, NOS
- 753.880 Other specified anomalies of bladder and urethra

753.9 Unspecified anomalies of urinary system

- 753.900 Unspecified anomaly of kidney
- 753.910 Unspecified anomaly of ureter
- 753.920 Unspecified anomaly of bladder
- 753.930 Unspecified anomaly of urethra
- 753.990 Unspecified anomaly of urinary system, NOS

754 Certain Congenital Musculoskeletal Anomalies

754.0 Of skull, face, and jaw Excludes: dentofacial anomalies (524.0) Pierre Robin sequence (524.080) syphilitic saddle nose (090.000) 754.000 Asymmetry of face 754.010 Compression (Potter's) facies # 754.020 Congenital deviation of nasal septum bent nose Τ 754.030 Dolichocephaly Always code if ≥36 weeks gestation # If <36 weeks gestation, code only if another reportable defect is present 754.040 Depressions in skull Includes: large fontanelle small fontanelle 754.050 Plagiocephaly 754.055 Asymmetric head T 754.060 Scaphocephaly, no mention of craniosynostosis * 754.070 Trigonocephaly, no mention of craniosynostosis Always code if ≥ 36 weeks gestation # If <36 weeks gestation, code only if another reportable defect is present * 754.080 Other specified skull deformity, no mention of craniosynostosis Includes:brachycephaly acrocephaly turricephaly oxycephaly

754.1 Anomalies of sternocleidomastoid muscle

Deformity of skull, NOS

* 754.090

754.100 Anomalies of sternocleidomastoid muscle

* Includes: absent or hypoplastic sternocleidomastoid contracture of sternocleidomastoid muscle sternomastoid tumor

Excludes: congenital sternocleidomastoid torticollis (use 756.860)

754.2 Certain congenital musculoskeletal deformities of spine

754.200 Congenital postural scoliosis 754.210 Congenital postural lordosis 754.220 Congenital postural curvature of spine, NOS

754.3 Congenital dislocation of hip

754.300 Congenital dislocation of hip
754.310 Unstable hip
preluxation of hip
subluxation of hip
predislocation status of hip at birth

754.4 Congenital genu recurvatum and bowing of long bones of leg

- 754.400 Bowing, femur
- 754.410 Bowing, tibia and/or fibula
- 754.420 Bow legs, NOS
- 754.430 Genu recurvatum
- 754.440 Dislocation of knee, congenital
- 754.490 Deformity of leg, NOS

754.5 Varus (inward) deformities of feet

- 754.500 Talipes equinovarus 754.510 Talipes calcaneovarus
- # 754.520 Metatarsus varus or metatarsus adductus
 - 754.530 Complex varus deformities
 - 754.590 Unspecified varus deformities of feet

754.6 Valgus (outward) deformities of feet

- 754.600 Talipes calcaneovalgus
- 754.610 Congenital pes planus
- 754.615 Pes valgus
- 754.680 Other specified valgus deformities of foot
- 754.690 Unspecified valgus deformities of foot

754.7 Other deformities of feet

- 754.700 Pes cavus
 - Claw foot (use 755.350 for claw foot)
- 754.720 Short Achilles tendon
- 754.730 Clubfoot, NOS
 - talipes, NOS
- 754.735 Congenital deformities of foot, NOS
- 754.780 Other specified deformities of ankle and/or toes
 - Includes: dorsiflexion of foot
 - Excludes: widely spaced 1st and 2nd toes (use 755.600)

754.8 Other specified congenital musculoskeletal deformities

- 754.800 Pigeon chest (pectus carinatum)
- 754.810 Funnel chest (pectus excavatum)
- 754.820 Other anomalies of chest wall
 - Includes: deformed chest, barrel chest
- 754.825 Shield chest
- 754.830 Dislocation of elbow
- 754.840 Club hand or fingers
- 754.850 Spade-like hand
- 754.880 Other specified deformity of hands
 - (see 755.500 for specified anomalies of fingers)

755 Other Congenital Anomalies of Limbs

755.0 Polydactyly

- 755.005 Accessory fingers (postaxial polydactyly, Type A) # 755.006 Skin tag (postaxial polydactyly, Type B) Exclude: Do not code in black infants. 755.007 Unspecified finger or skin tag (postaxial polydactyly, NOS)
 755.010 Accessory thumbs (preaxial polydactyly)
 755.020 Accessory toes (postaxial)
 755.030 Accessory big toe (preaxial)
 755.090 Accessory digits, NOS (hand/foot not specified)
 755.095 Accessory digits hand, NOS (preaxial, postaxial not
 - specified)
 - 755.096 Accessory digits foot, NOS (preaxial, postaxial not specified)

755.1 Syndactyly

- 755.100 Fused fingers
- 755.110 Webbed fingers
- 755.120 Fused toes
- # 755.130 Webbed toes

Code webbing of the second and third toes only if another reportable defect is present. Always code webbing of other toes regardless of whether another reportable defect is present

- 755.190 Unspecified syndactyly (see below for specified site)
- 755.191 Unspecified syndactyly thumb and/or fingers, unilateral
- 755.192 Unspecified syndactyly thumb and/or fingers, bilateral
- 755.193 Unspecified (webbed vs. fused) syndactyly thumb and/or fingers, NOS
- 755.194 Unspecified syndactyly toes unilateral
- 755.195 Unspecified syndactyly toes bilateral
- 755.196 Unspecified syndactyly toes, NOS
- 755.199 Unspecified syndactyly (i.e., webbed vs. fused) digits not known

755.2 Reduction defects of upper limb

- Т If description of the condition includes amniotic or constricting bands use additional code, 658.800 (Only use 658.800 if another reportable defect is present) Excludes shortening of upper limb (use 755.580) or hypoplasia of upper limb (use 755.585)
 - 755.200 Absence of upper limb

humerus (total or partial), radius, ulna and hand Absent:

Includes: amelia of upper limb, NOS

infants with rudimentary or nubbin fingers

attached to stump of humerus or shoulder girdle

755.210 Absence of upper arm and forearm

humerus (total or partial), radius and ulna Absent:

(total or partial)

Present: hand (total or partial)

Includes: phocomelia of upper limb, NOS;

intercalary reduction defect of upper limb, NOS

755.220 Absence of forearm only or upper arm only Absent: radius and ulna Present: humerus, hand (total or partial) or Absent: humerus Present: radius, ulna, and hand 755.230 Absence of forearm and hand Absent: radius and ulna (total or partial) and hand Includes: infants with rudimentary or nubbin fingers attached to stump of forearm or elbow 755.240 Absence of hand or fingers hand or fingers (total or partial) not in Absent: conjunction with ray or long bone reduction Includes: rudimentary or nubbin fingers; absent individual phalanges; absent or missing fingers, NOS Excludes: isolated absent or hypoplastic thumb (use 755.260) 755.250 Split-hand malformation central fingers (third with or without second, Absent: fourth) and metacarpals (total or partial) Includes: monodactyly; lobster-claw hand Excludes: isolated absent central fingers without metacarpal defects (use 755.240) 755.260 Preaxial longitudinal reduction defect of upper limb radius (total or partial) and/or thumb with or Absent: without second finger (total or partial) Includes: isolated absent or hypoplastic thumb; radial ray defect, NOS 755.265 Longitudinal reduction defect of upper limb, NOS Includes: absent forearm long bone with absent fingers, 755.270 Postaxial longitudinal reduction defect of upper limb Includes: isolated absent ulna (total or partial); absent fifth with or without fourth finger (total or partial) only if ulna or fifth \pm fourth metacarpal also totally or partially absent; ulnar ray defect, NOS 755.280 Other specified reduction defect of upper limb 755.285 Transverse reduction defect of upper limb, NOS Includes: congenital amputation of upper limb, NOS

755.3 Reduction defects of lower limb

T If description of condition includes amniotic or constricting bands use additional code, 658.800 (Only use this code if another reportable defect is present)

755.290 Unspecified reduction defect of upper limb

Excludes shortening of lower limb (use 755.680) and hypoplasia of lower limb (use 755.685)

755.300 Absence of lower limb femur (total or partial), tibia, fibula, and Absent: Includes: amelia of lower limb, NOS infants with rudimentary or nubbin toes attached to stump of femur or pelvic girdle 755.310 Absence of thigh and lower leg Absent: femur (total or partial), tibia and fibula (total or partial) Present: foot (total or partial) Includes: phocomelia of lower limb, NOS; intercalary reduction defect of lower limb, NOS 755.320 Absence of lower leg only or femur only Absent: tibia and fibula Present: femur, foot (total or partial) orAbsent: femur Present: tibia, fibula, and foot 755.330 Absence of lower leg and foot Absent: tibia and fibula (total or partial), foot Includes: infants with rudimentary or nubbin toes attached to stump of leg or knee 755.340 Absence of foot or toes Absent: foot or toes (total or partial) not in conjunction with ray or long bone reduction Includes: rudimentary or nubbin toes; absent individual phalanges; absent or missing toes, NOS Excludes: isolated absent or hypoplastic great toe (use 755.365) Split-foot malformation 755.350 central toes (third with or without second, Absent: fourth) and metatarsals (total or partial) Includes: monodactyly; lobster claw foot Excludes: isolated absent central toes without metatarsal defects (use 755.340) Note: preaxial lower limb reductions can occur with split-hand malformations of the upper limb and these lower limb defects should be coded 755.365 755.360 Longitudinal reduction defect of lower limb, NOS Includes: absent long bone of leg with absent toes, NOS 755.365 Preaxial longitudinal reduction defect of lower limb tibia (total or partial) and/OR great toe with Absent: or without second toe (total or partial) Includes: isolated absent or hypoplastic great toe; tibial ray defect, NOS 755.366 Postaxial longitudinal reduction defect of lower limb Includes: isolated absent fibula (total or partial); absent fifth with or without fourth toe (total or partial) only if fibula or fifth ± fourth metatarsal also totally or partially absent; fibular ray defect, NOS 755.380 Other specified reduction defect of lower limb 755.385 Transverse reduction defect of lower limb, NOS Includes: congenital amputation of lower limb, NOS 755.390 Unspecified reduction defect of lower limb

755.4 Reduction defects of unspecified limb

- T If description of condition includes amniotic or constricting bands use additional code, 658.800 (note: 658.00 should only be used with another reportable defect)
 - 755.400 Absence of limb, NOS

Includes: amelia, NOS

755.410 Phocomelia, NOS

Includes: intercalary reduction defect, NOS

755.420 Transverse reduction defect, NOS

Includes: congenital amputation of unspecified limb

755.430 Longitudinal reduction defect, NOS

Includes: preaxial or postaxial reduction defect, NOS

755.440 Absent digits, not specified whether fingers or toes

755.480 Other specified reduction defect of unspecified limb

755.490 Unspecified reduction defect of unspecified limb

755.5 Other anomalies of upper limb, including shoulder girdle

Includes: complex anomalies involving all or part of upper limb

755.500 Anomalies of fingers

Includes: camptodactyly

> clinodactyly macrodactylia brachydactyly

triphalangeal thumb incurving fingers

Excludes: acrocephalosyndactyly (see 756.050)

Apert's syndrome (see 756.055)

755.510 Anomalies of hand

Excludes: simian crease (use 757.200)

- 755.520 Anomalies of wrist 755.525 Accessory carpal bones
- 755.526 Madelung's deformity
- 755.530 Anomalies of forearm, NOS
- 755.535 Radioulnar dysostosis
- 755.536 Radioulnar synostosis
- 755.540 Anomalies of elbow and upper arm
- 755.550 Anomalies of shoulder
- 755.555 Cleidocranial dysostosis 755.556 Sprengel's deformity
- 755.560 Other anomalies of whole arm
- 755.580 Other specified anomalies of upper limb

Includes: hyperextensibility of upper limb

shortening of arm

755.585 Hypoplasia of upper limb

Includes: hypoplasia of fingers, hands, or arms

Excludes: aplasia or absent upper limb (see 755.2)

755.590 Unspecified anomalies of upper limb

755.6 Other anomalies of lower limb, including pelvic girdle

Includes:complex anomalies involving all
 or part of lower limb

755.600 Anomalies of toes

Includes: overlapping toes

hammer toes

widely spaced first and second toes

755.605 Hallux valgus

755.606 Hallux varus

755.610 Anomalies of foot

Includes: plantar furrow

Excludes: lobster claw foot (use 755.350)

755.616 Rocker-bottom foot

755.620 Anomalies of ankle

astragaloscaphoid synostosis

755.630 Anomalies of lower leg

angulation of tibia, tibial torsion

(exclude if clubfoot present)

755.640 Anomalies of knee

hyperextended knee

755.645 Genu valgum

755.646 Genu varum

755.647 Absent patella or rudimentary patella

755.650 Anomalies of upper leg

anteversion of femur

755.660 Anomalies of hip

Includes: coxa vara

coxa valga

other abnormalities of hips

755.665 Hip dysplasia, NOS

755.666 Unilateral hip dysplasia

755.667 Bilateral hip dysplasia

755.670 Anomalies of pelvis

fusion of sacroiliac joint

755.680 Other specified anomalies of lower limb

hyperextended legs

shortening of legs

755.685 Hypoplasia of lower limb

Includes: hypoplasia of toes, feet, legs

Excludes: aplasia or absent lower limb (see 755.3)

755.690 Unspecified anomalies of legs

755.8 Other specified anomalies of unspecified limb

755.800 Arthrogryposis multiplex congenita

Includes: distal arthrogryposis syndrome

755.810 Larsen's syndrome

Includes:

755.880 Other specified anomalies of unspecified limb

overlapping digits, NOS

hyperextended joints, NOS

Excludes: hyperextended knees (use 755.640)

755.9 Unspecified anomalies of unspecified limb

S = Rev. 8/07 R = Rev. 6/07 N = Rev. 5/07

T = Rev. 6/04

I = Rev. 6/04

* = code created by CDC

= on the MACDP Excl List

756 Other Congenital Musculoskeletal Anomalies

756.0 Anomalies of skull and face bones

Excludes: skull and face deformities in 754 Pierre Robin sequence (use 524.080) 756.000 Craniosynostosis, NOS craniostenosis, NOS closed-skull sutures, NOS 756.005 Sagittal craniosynostosis 756.006 Metopic craniosynostosis 756.010 Coronal craniosynostosis 756.020 Lambdoidal craniosynostosis 756.030 Other types of craniosynostosis Includes: basilar craniosynostosis 756.040 Craniofacial dysostosis Includes: Crouzon's disease 756.045 Mandibulofacial dysostosis Includes: Franceschetti syndrome Treacher-Collins syndrome 756.046 Other craniofacial syndromes Includes: oculomandibulofacial syndrome Hallermann-Streiff syndrome 756.050 Acrocephalosyndactyly, NOS 756.055 Acrocephalosyndactyly types I or II Apert syndrome 756.056 Acrocephalosyndactyly type III 756.057 Other specified acrocephalosyndactylies 756.060 Goldenhar syndrome oculoauriculovertebral dysplasia 756.065 Hemifacial microsomia 756.080 Other specified skull and face bone anomalies localized skull defects Includes: # flat occiput mid-facial hypoplasia # prominent occiput prominent maxilla hypotelorism Excludes: macrocephaly (use 742.400) small chin (see 524.0) Pierre Robin sequence (use 524.080) 756.085 Hypertelorism, telecanthus, wide set eyes 756.090 Unspecified skull and face bone anomalies Excludes: dentofacial anomalies (524.0) skull defects associated with brain anomalies such as: anencephalus (740.0) encephalocele (742.0) hydrocephalus (742.3) microcephalus (742.100)

756.1 Anomalies of spine

756.100 Spina bifida occulta 756.110 Klippel-Feil syndrome Wildervanck syndrome

| 756.120 | Kyphosis kyphoscoliosis |
|---------|---|
| 756.130 | Congenital spondylolisthesis |
| 756.140 | Anomalies of cervical vertebrae |
| 756.145 | Hemivertebrae (cervical) |
| 756.146 | Agenesis (cervical) |
| 756.150 | Anomalies of thoracic vertebrae |
| | Hemivertebrae of thoracic vertebrae |
| 756.156 | Agenesis of thoracic vertebrae |
| 756.160 | Anomalies of lumbar vertebrae |
| | Hemivertebrae of lumbar vertebrae |
| 756.166 | Agenesis of lumbar vertebrae |
| 756.170 | Sacrococcygeal anomalies |
| | Includes: agenesis of sacrum |
| | Excludes: pilonidal sinus (see 685.100) |
| 756.179 | Sacral mass, NOS |
| 756.180 | Other specified vertebral anomalies |
| 756.185 | Hemivertebrae, NOS |
| 756.190 | Unspecified anomalies of spine |

756.2 Cervical rib

756.200 Cervical rib supernumerary rib in cervical region

756.3 Other anomalies of ribs and sternum

| 756.300 | Absence of ribs | | | | | |
|---------|--|--|--|--|--|--|
| 756.310 | Misshapen ribs | | | | | |
| 756.320 | Fused ribs | | | | | |
| 756.330 | Extra ribs | | | | | |
| 756.340 | Other anomalies of ribs | | | | | |
| 756.350 | Absence of sternum | | | | | |
| 756.360 | Misshapen sternum | | | | | |
| 756.380 | Other anomalies of sternum | | | | | |
| | Includes: double ossification center in the manubrium, | | | | | |
| | bifid sternum, short sternum | | | | | |
| 756.390 | Anomalies of thoracic cage, unspecified | | | | | |
| | Excludes: deformed chest (use 754.820) | | | | | |

756.4 Chondrodystrophy

| 756.400 | Asphyxiating thoracic dystrophy Jeune syndrome |
|---------|--|
| | thoracic-pelvic-phalangeal dysplasia |
| | Excludes: homozygous achondroplasia |
| 756.410 | Chondrodysplasia |
| | Ollier syndrome, enchondromatosis |
| 756.420 | Chondrodysplasia with hemangioma |
| | Kast syndrome |
| | Maffucci syndrome |
| 756.430 | Achondroplastic dwarfism |
| 756.440 | Other specified chondrodystrophies |
| | Excludes: Conradi's (use 756.575) |
| 756.445 | Diastrophic dwarfism |
| 756.446 | Metatrophic dwarfism |
| 756.447 | Thanatophoric dwarfism |
| / | I |

```
756.450 Metaphyseal dysostosis
      756.460 Spondyloepiphyseal dysplasia
      756.470 Exostosis
               Excludes:
                             Gardner syndrome (see 759.630)
      756.480 Other specified chondrodystrophy
      756.490 Unspecified chondrodystrophy
               Excludes:
                             lipochondrodystrophy (use 277.510)
756.5 Osteodystrophies
      756.500 Osteogenesis imperfecta
      756.505 Osteopsathyrosis
      756.506 Fragilitas ossium
      756.510 Polyostotic fibrous dysplasia
               Albright-McCune-Sternberg syndrome
      756.520 Chondroectodermal dysplasia
      756.525 Ellis-van Creveld syndrome
      756.530 Infantile cortical hyperostosis
               Caffey syndrome
      756.540 Osteopetrosis
               Albers-Schonberg syndrome
               marble bones
      756.550 Progressive diaphyseal dysplasia
               Engelmann syndrome
               Camurati-Engelmann disease
      756.560 Osteopoikilosis
756.570 Multiple epiphyseal dysplasia
      756.575 Conradi syndrome
               chondrodysplasia punctata
               Excludes:
                            warfarin embryopathy
      756.580 Other specified osteodystrophies
      756.590 Unspecified osteodystrophies
756.6 Anomalies of diaphragm
      756.600 Absence of diaphragm
      756.610 Congenital diaphragmatic hernia
      756.615 Diaphragmatic hernia (Bochdalek)
      756.616 Diaphragmatic hernia (Morgagni)
      756.617 Hemidiaphragm
      756.620 Eventration of diaphragm
      756.680 Other specified anomalies of diaphragm
      756.690 Unspecified anomalies of diaphragm
      756.700 Exomphalos, omphalocele
      756.710 Gastroschisis
               Excludes:
                             umbilical hernia (553.100)
      756.720 Prune belly syndrome
```

756.7 Anomalies of abdominal wall

```
756.790 Other and unspecified anomalies of abdominal wall
756.795 Epigastric hernia
```

756.8 Other specified anomalies of muscle, tendon, fascia and connective tissue

```
756.800 Poland syndrome or anomaly
756.810 Other absent or hypoplastic muscle
```

Includes: absent pectoralis major
Excludes: prune belly syndrome (use 756.720)

756.820 Absent tendon

756.830 Nail-patella syndrome
756.840 Amyotrophia congenita

756.850 Ehlers-Danlos syndrome
756.860 Congenital torticollis
(see also 754.100, anomalies of sternocleidomastoid muscle)

756.880 Other specified anomalies of muscle, tendon, fascia and connective tissue
Includes: myopathy, congenital NOS

756.9 Unspecified anomalies of musculoskeletal system

| 756.900 | Unspecified | anomalies | of | muscle |
|---------|-------------|-----------|----|------------------------|
| 756.910 | Unspecified | anomalies | of | tendon |
| 756.920 | Unspecified | anomalies | of | bone |
| 756.930 | Unspecified | anomalies | of | cartilage |
| 756.940 | Unspecified | anomalies | of | connective tissue |
| 756.990 | Unspecified | anomalies | of | musculoskeletal system |

757 Congenital Anomalies of the Integument

757.000 Hereditary edema of legs Hereditary trophedema Milroy's disease

757.1 Ichthyosis congenita

757.100 Harlequin fetus
757.110 Collodion baby
757.115 Bullous type
757.120 Sjogren-Larsson syndrome
757.190 Other and unspecified
757.195 Ichthyosis vulgaris
757.196 X-linked ichthyosis
757.197 Ichthyosiform erythroderma

757.2 Dermatoglyphic anomalies

757.200 Abnormal palmar creases
Includes: simian creases, transverse palmar creases

757.3 Other specified anomalies of skin

Excludes:pigmented mole (216.900) hemangioma (see 228.0)

757.300 Specified syndromes, not elsewhere classified, involving skin anomalies

757.310 Skin tags

Includes: anal tags

Excludes: preauricular tag (see 744.110) vaginal tags (see 752.480)

757.320 Urticaria pigmentosa 757.330 Epidermolysis bullosa

757.340 Ectodermal dysplasia

Excludes: Ellis-van Creveld syndrome (756.525)

757.345 X-linked type ectodermal dysplasia 757.346 Other specified ectodermal dysplasias

757.350 Incontinentia pigmenti 757.360 Xeroderma pigmentosum

757.370 Cutis laxa hyperelastica

757.380 Nevus, not elsewhere classifiable

Includes: port wine stain or nevus flammeus

T Excludes: hairy nevus (use 216.920)

Sturge-Weber syndrome (use 759.610)

757.385 Birthmark, NOS

757.386 Mongolian blue spot

757.390 Other specified anomalies of skin

Includes: cafe au lait spots hyperpigmented areas

skin cysts

hypoplastic dermal patterns

757.395 Absence of skin

757.4 Specified anomalies of hair

Excludes: kinky hair syndrome (use 759.870)

757.400 Congenital alopecia

Excludes: ectodermal dysplasia (use 757.340)

757.410 Beaded hair

Monilethrix

757.420 Twisted hair

Pili torti

757.430 Taenzer's hair

757.450 Persistent or excessive lanugo

Includes: hirsutism

757.480 Other specified anomalies of hair

757.5 Specified anomalies of nails

757.500 Congenital anonychia

Absent nails

757.510 Enlarged or hypertrophic nails

757.515 Onychauxis 757.516 Pachyonychia

757.520 Congenital koilonychia

757.530 Congenital leukonychia

757.540 Club nail

757.580 Other specified anomalies of nails

757.585 Hypoplastic (small) fingernails and/or toenails

757.6 Specified anomalies of breast

757.600 Absent breast with absent nipple

757.610 Hypoplastic breast with hypoplastic nipple

757.620 Accessory (ectopic) breast with nipple

757.630 Absent nipple

Т # 757.640 Small nipple (hypoplastic)

Always code if ≥36 weeks gestation

If <36 weeks gestation, code only if another reportable defect is present

757.650 Accessory (ectopic) nipple, supernumerary

757.680 Other specified anomalies of breast

Widely spaced nipples

Excludes: inverted nipples (never a defect)

757.8 Other specified anomalies of the integument

757.800 Includes: scalp defects

For specified anomalies of skin see 757.390 For specified anomalies of hair see 757.480

For specified anomalies of nails see 757.580

757.9 Unspecified anomalies of the integument

757.900 Unspecified anomalies of skin

757.910 Unspecified anomalies of hair, NOS

757.920 Unspecified anomalies of nail, NOS

757.990 Unspecified anomalies of the integument, NOS

S = Rev. 8/07R = Rev. 6/07N = Rev. 5/07T = Rev. 6/04* = code created by CDC

= on the MACDP Excl List

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758 Chromosomal Anomalies

758.0 Down syndrome

Clinical Down syndrome karyotype identified as:

- Т 758.000 Down syndrome, karyotype trisomy 21, cytogenetics result in record
- Т 758.008 Down syndrome suspected, cytogenetics pending
 - 758.010 Down syndrome, karyotype trisomy G, NOS

 - 758.020 Translocation trisomy duplication of a 21 758.030 Translocation trisomy duplication of a G, NOS
 - 758.040 Mosaic Down syndrome
- Т 758.090 Down syndrome, NOS (i.e. chart states a diagnosis of Trisomy 21 or Downs syndrome, but no cytogenetics result in record)
- Т 758.098 Down syndrome suspected, cytogenetics never done

758.1 Patau syndrome

Clinical Patau syndrome karyotype identified as:

- Т 758.100 Patau syndrome, karyotype trisomy 13, cytogenetics result in record
 - 758.108 Patau syndrome suspected, cytogenetics pending
 - 758.110 Patau syndrome, karyotype trisomy D, NOS
 - 758.120 Translocation trisomy duplication of a 13
- 758.130 Translocation trisomy duplication of a D, NOS
- Т 758.190 Patau syndrome, NOS (i.e. chart states a diagnosis of Trisomy 13 or Patau syndrome, but no cytogenetics result in record)
- Т 758.198 Patau syndrome suspected, cytogenetics pending

758.2 Edwards syndrome

Clinical Edwards syndrome karyotype identified as:

- Т 758.200 Edwards syndrome, karyotype trisomy 18, cytogenetics result in record
- Т 758.208 Edwards syndrome suspected, cytogenetics pending
 - 758.210 Edwards syndrome, karyotype trisomy E, NOS

 - 758.220 Translocation trisomy duplication of an 18 758.230 Translocation trisomy duplication of an E, NOS
- 758.290 Edwards syndrome, NOS (i.e. chart states a diagnosis of Т Trisomy 18 or Edwards syndrome, but no cytogenetics
 - result in record) 758.295 Edwards phenotype - normal karyotype
- Т 758.298 Edwards syndrome suspected, cytogenetics pending

758.3 Autosomal deletion syndromes

```
758.300 Antimongolism syndrome
         Clinical antimongolism syndrome:
           karyotype - partial or total deletion of:
                 21
                 G, NOS
           NOS
758.310 Cri du chat syndrome
         Clinical Cri du chat syndrome:
           karyotype - deletion of:
                 5
                 B, NOS
           NOS
758.320 Wolff-Hirschorn syndrome
         Clinical Wolff-Hirschorn syndrome:
           karyotype - deletion of:
                 4
                 B, NOS
           NOS
758.330 Deletion of long arm of 13
deletion of long arm of D, NOS 758.340 Deletion of long arm of E \,
         deletion of long arm of 17 or 18
758.350 Deletion of short arm of E
         deletion of short arm of 17 or 18
758.360 Monosomy G mosaicism
758.370 Deletion in band 11 of long arm of 22 (22g11 deletions)
         Note: Code added for use with births on or after 4/1/2001
758.380 Other loss of autosomal material
758.390 Unspecified autosomal deletion syndromes
```

758.4 Balanced autosomal translocation in normal individual

758.400 Balanced autosomal translocation in normal individual

758.5 Other conditions due to autosomal anomalies

```
758.500
        Trisomy 8
758.510
        Other trisomy C syndromes
        Trisomy: 6, 7, 9, 10, 11, 12, or C, NOS
758.520 Other total trisomy syndromes
        Trisomy 22
        Trisomy, NOS
758.530 Partial trisomy syndromes
758.540 Other translocations
                     balanced translocation in normal
        Excludes:
                      individual (use 758.400)
758.580 Other specified anomalies of autosomes, NOS
        Includes:
                      marker autosome
758.585 Polyploidy
758.586 Triploidy
758.590 Unspecified anomalies of autosomes
```

758.6 Gonadal Dysgenesis

Excludes:pure gonadal dysgenesis (752.720) Noonan syndrome (759.800)

758.600 Turner's phenotype, karyotype 45, X [XO]

758.610 Turner's phenotype, variant karyotypes

karyotype characterized by:

isochromosome

mosaic, including XO

partial X deletion

ring chromosome

Excludes: Turner's phenotype, karyotype normal XX

(use 759.800, Noonan syndrome)

758.690 Turner syndrome, karyotype unspecified, NOS

Bonneville-Ullrich syndrome, NOS

758.7 Klinefelter syndrome

758.700 Klinefelter's phenotype, karyotype 47, XXY

758.710 Klinefelter's phenotype, other karyotype with additional

X chromosomes

XX

XXXY

XXYY

XXXXY

758.790 Klinefelter syndrome, NOS

758.8 Other conditions due to sex chromosome anomalies

758.800 Mosaic XO/XY, 45X/46XY

Excludes: with Turner's phenotype (758.610)

758.810 Mosaic XO/XX

Excludes: with Turner's phenotype (758.610)

758.820 Mosaic XY/XXY,46XY/47XXY

Excludes: Klinefelter's phenotype (758.710)

758.830 Mosaic including XXXXY, 49XXXXY

Excludes: with Klinefelter's phenotype (use 758.710)

758.840 XYY, male, 47XYY

mosaic XYY male

758.850 XXX female, 47XXX

758.860 Additional sex chromosomes, NOS

758.880 Other specified sex chromosome anomaly

Includes: fragile X

758.890 Unspecified sex chromosome anomaly

758.9 Conditions due to anomaly of unspecified chromosomes

758.900 Mosaicism, NOS

758.910 Additional chromosome(s), NOS

758.920 Deletion of chromosome(s), NOS

758.930 Duplication of chromosome(s), NOS

758.990 Unspecified anomaly of chromosome(s)

759 Other and Unspecified Congenital Anomalies

759.0 Anomalies of spleen

```
759.000 Absence of spleen
asplenia
759.005 Ivemark syndrome
759.010 Hypoplasia of spleen
# 759.020 Hyperplasia of spleen
splenomegaly
hepatosplenomegaly (also use code 751.620)
759.030 Misshapen spleen
759.040 Accessory spleen
759.050 Ectopic spleen
759.080 Other specified anomalies of spleen
759.090 Unspecified anomalies of spleen
```

759.1 Anomalies of adrenal gland

| 759.100 | Absence of adrenal gland |
|---------|--|
| 759.110 | Hypoplasia of adrenal gland |
| 759.120 | Accessory adrenal gland |
| 759.130 | Ectopic adrenal gland |
| 759.180 | Other specified anomaly of adrenal gland |
| | Excludes: congenital adrenal hyperplasia |
| | (use 255.200) |
| 759.190 | Unspecified anomalies of adrenal gland |

759.2 Anomalies of other endocrine glands

| | 759.200 | Anomalies of pituitary gland |
|---|---------|--|
| | 759.210 | Anomalies of thyroid gland |
| | 759.220 | Thyroglossal duct anomalies |
| | | thyroglossal cyst |
| | 759.230 | Anomalies of parathyroid gland |
| # | 759.240 | Anomalies of thymus |
| | | thymic hypertrophy |
| | | absent thymus |
| | 759.280 | Other specified anomalies of endocrine gland |
| | 759.290 | Unspecified anomaly of endocrine gland |

759.3 Situs inversus

| 759.300 | Dextrocardia with complete situs inversus |
|---------|---|
| 759.310 | Situs inversus with levocardia |
| 759.320 | Situs inversus thoracis |
| 759.330 | Situs inversus abdominis |
| 759.340 | Kartagener syndrome (triad) |
| 759.390 | Unspecified situs inversus |
| | Excludes: dextrocardia (746.800) not |
| | associated with complete situs inversus |

759.4 Conjoined twins

```
759.400 Dicephalus
two heads
759.410 Craniopagus
```

head-joined twins
759.420 Thoracopagus
thorax-joined twins
759.430 Xiphopagus
xiphoid- and pelvis-joined twins
759.440 Pygopagus
buttock-joined twins
759.480 Other specified conjoined twins

759.5 Tuberous sclerosis

759.500 Tuberous sclerosis
Bourneville's disease
epiloia

759.6 Other hamartoses, not elsewhere classified

759.490 Unspecified conjoined twins

759.600 Peutz-Jeghers syndrome
759.610 Encephalocutaneous angiomatosis
Kalischer's disease
Sturge-Weber syndrome
759.620 Von Hippel-Lindau syndrome
759.630 Gardner syndrome
759.680 Other specified hamartomas
759.690 Unspecified hamartomas

759.7 Multiple congenital anomalies,

759.700 Multiple congenital anomalies, anomaly, multiple, NOS deformity, multiple, NOS

759.8 Other specified anomalies and syndromes

cyclops Noonan syndrome oral-facial-digital (OFD) syndrome, type I Orofaciodigital syndrome, type II (Mohr syndrome) Waardenburg syndrome whistling face syndrome 759.820 Cong malformation syndromes associated with short stature Amsterdam dwarf (Cornelia de Lange syndrome) Cockayne syndrome Laurence-Moon-Biedl syndrome Russell-Silver syndrome Seckel syndrome Smith-Lemli-Opitz syndrome 759.840 Cong malformation syndromes involving limbs Carpenter syndrome Holt-Oram syndrome Klippel-Trenaunay-Weber syndrome Rubinstein-Taybi syndrome sirenomelia thrombocytopenia-absent radius (TAR) syndrome

759.860 Cong malformation syndromes with other skeletal changes

759.800 Cong malformation syndromes affecting facial appearance

Marfan syndrome Stickler syndrome

759.870 Cong malformation syndromes with metabolic disturbances

Alport syndrome

Beckwith (Wiedemann-Beckwith) syndrome

leprechaunism

Menkes syndrome (kinky hair syndrome)

Prader-Willi syndrome Zellweger syndrome

759.890 Other specified anomalies

Includes: hemihypertrophy

Meckel-Gruber syndrome

759.9 Congenital anomaly, unspecified

759.900 Anomalies of umbilicus

low-lying umbilicus umbilical cord atrophy

759.910 Embryopathia, NEC 759.990 Congenital anomaly, NOS

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List ordered alphabetically
       524.000 Abnormalities of jaw size
               micrognathia
               macrognathia T
       255.200
              Adrenogenital syndrome
      270.200 Albinism
    # 277.620 Alpha-1 antitrypsin deficiency
    # 658.800 Amniotic bands (constricting bands, amniotic cyst)
    # 270.600 Arginosuccinic aciduria
    # 778.000 Ascites, congenital
216
      Benign neoplasm of skin
    T (NOTE: All neoplasms should be coded ONLY if another reportable code
        is present)
                Includes: blue nevus
                                          pigmented nevus
                          papilloma
                                          dermatofibroma
                          syringoadenoma hydrocystoma
                        * dermoid cyst
                                        syringoma
               Excludes: skin of female genital organs (use 221.000),
                          skin of male genital organs (use 222.000)
T
               Benign neoplasm of skin, ear and external auditory canal
    # 216.200
                Includes: auricle ear
                           external meatus
                           auricular canal
                           external canal
                           pinna
                             cartilage of ear
               Excludes:
Т
    # 216.100
               Benign neoplasm of skin, eyelid, including canthus
               Excludes:
                             cartilage of eyelid
Τ
    # 216.000
               Benign neoplasm of skin, lip
               Excludes:
                             vermillion border of lip
    # 216.700 Benign neoplasm of skin, lower limb, hip
Т
    # 216.300
               Benign neoplasm of skin, other and unspecified parts of face
                Includes:
                             cheek, external nose, external eyebrow
                             temple
Т
    # 216.800
              Benign neoplasm of skin, other specified sites of skin
               Excludes:
                             epibulbar dermoid cyst (use 743.810)
T
    # 216.400
               Benign neoplasm of skin, scalp and skin of neck
Т
    # 216.900
               Benign neoplasm of skin, site unspecified
               Benign neoplasm of skin, trunk, except scrotum
    # 216.500
                             axillary fold
                Includes:
                             perianal skin
                             skin of:
                                                 wall,
                                                         abdominal wall,
                                          chest
                                          buttock, anus, perineum, back, umbilicus,
                                          breast
                             anal canal
               Excludes:
                             anus, NOS
                             skin of scrotum
```

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T
      216.600
               Benign neoplasm of skin, upper limb, shoulder
               Benign skin neoplasm of female genital organs
      221.000
      222.000
               Benign skin neoplasm of male genital organs
      453.000 Budd-Chiari, occlusion of hepatic vein
      427.900 Cardiac arrhythmias, NEC. Never code premature atrial
               contractions, PACs.
     330.100
              Cerebral lipidoses
                            Tay-Sachs disease, gangliosidosis
               Includes:
      363.200
               Chorioretinitis
      279.200
               Combined immunodeficiency syndrome
      771.280
               Congenital infection, other specified
                             human immunodeficiency virus (HIV)
               Excludes:
                             infection and acquired immunodeficiency
                             syndrome (AIDS)
    # 277.000 Cystic fibrosis
               No mention of meconium ileus
     277.010 Cystic fibrosis
               With mention of meconium ileus
      228.100 Cystic hygroma
               Lymphangioma, any site
      771.100 Cytomegalovirus (CMV) (in utero infections only)
      253.820
               Diencephalic syndrome
      279.110
               DiGeorge syndrome
      277.400 Disorders of bilirubin excretion
      425.300 Endocardial fibroelastosis
      553.200
               Epigastric hernia
      767.600
               Erb's palsy
      368.000
               Esotropia
      378.000
               Exotropia
      351.000
               Facial palsy
      331.890 Familial degenerative CNS disease
      760.710 Fetal alcohol syndrome
      760.718 Fetal alcohol syndrome, probable
               Includes: "facies"
      760.750 Fetal hydantoin (Dilantin) syndrome
      282.200 Glucose-6-phosphate dehydrogenase (G-6-PD) deficiency
      271.000 Glycogen storage diseases
Τ
      216.920 Hairy nevus
228.0 Hemangioma
                Include if greater than 4-inches diameter, if multiple
               hemangiomas, or if cavernous hemangioma
      228.040
               Hemangioma, intra-abdominal (Always code regardless of size, type or
               number)
               Hemangioma, intracranial (Always code regardless of size, type or
      228.020
      228.090
               Hemangioma, of other sites (Always code regardless of size, type or
               number)
     228.000
               Hemangioma, of unspecified site. Always code if multiple hemangiomas
               of any size are present, if one or more cavernous hemangiomas of any
               size are present, or if a single hemangioma measuring \geq 4cm in
               diameter or desribed as large, huge, or of medical significance is
      228.030 Hemangioma, retinal (Always code regardless of size, type or number)
```

S = Rev. 8/07 R = Rev. 6/07 N = Rev. 5/07 T = Rev. 6/04 * = code created by CDC

= on the MACDP Excl List

Т

```
# 228.010
               Hemangioma, skin & subcutaneous, NOS Always code if multiple
               hemangiomas of any size are present, if one or more cavernous
               hemangiomas of any size are present, or if a single hemangioma
               measuring ≥ 4cm in diameter or desribed as large, huge, or of medical
               significance is present.
    # 286.000 Hemophilia (all types)
      774.490 Hepatitis, neonatal, NOS
      774.480 Hepatitis, neonatal, other specified
     282.100 Hereditary elliptocytosis
    # 282.000 Hereditary spherocytosis
      771.220 Herpes simplex (in utero infections only)
               Includes: encephalitis
                      meningoencephalitis
      202.300 Histiocytosis, malignant
      277.510 Hurler syndrome
               Includes:
                            lipochondrodystrophy
     778.600 Hydrocele, congenital
     270.700 Hyperglycinemia
     251.200 Hypoglycemia, idiopathic
     252.100 Hypoparathyroidism, congenital
     275.330 Hypophosphatemic rickets
      253.280 Hypopituitarism, congenital
              Hypothyroidism, congenital (Exclude even if other
    # 243.990
               defects are present only if the record specifies
               hypothyroidism of prematurity. Other types of
               hypothyroidism or hypothyroidism NOS should continue
               to be on the routine exclusion list.)
      345.600 Infantile spasms, congenital
    # 550.000 Inquinal hernia or patent processus vaginalis never
        -550.900 code in infants if <36 weeks gestation regardless of
                 the presence of a reportable defect.
                 NOTE: for those ≥36 weeks:
                 Code in males only if another reportable defect is present;
                 Code in females, always code even if found in isolation
      208.000 Leukemia, congenital, NOS
214
      Lipoma
      214.300 Lipoma, intra-abdominal organs
               Lipoma, intrathoracic organs
      214.200
      214.810 Lipoma, lumbar or sacral lipoma
                       paraspinal lipoma
      214.100 Lipoma, other skin and subcutaneous tissue
      214.800 Lipoma, other specified sites
      214.000 Lipoma, skin and subcutaneous tissue of face
      214.400 Lipoma, spermatic cord
      214.900 Lipoma, unspecified site
    # 457.800 Lymphatics - other specified disorders of (including chylothorax)
      524.000 Macrognathia
     270.300 Maple syrup urine disease
     777.600 Meconium peritonitis
     777.100 Meconium plug syndrome
      524.000 Micrognathia
      352.600 Moebius syndrome
```

```
# 520.600 Natal teeth
239.200 Neck cyst
774.490 Neonatal hepatitis, NOS
```

```
774.480 Neonatal hepatitis, other specified
      159.800 Neoplasms of the abdomen, other specified
      191.000 Neoplasms of the CNS
               Includes:
                            medulloblastoma, gliomas
      171.800 Neoplasms of the connective tissue
                           Ewing's sarcoma
               Includes:
                             fibrosarcoma
      155.000 Neoplasms of the liver
               Includes:
                            hepatoblastoma
               hemangio-epithelioma
      162.800 Neoplasms of the lung
      186.000 Neoplasms of the testes
      194.000 Neuroblastoma
      237.700 Neurofibromatosis
     379.500 Nystagmus
    # 270.100 Phenylketonuria (PKU)
    * 524.080 Pierre Robin sequence
    # 685.100 Pilonidal sinus (sacrodermal), sacral sinus, sacral dimple
    # 277.630 Pseudocholinesterase enzyme deficiency
    # 284.000 Red cell aplasia
      362.600 Retinal degeneration, peripheral
      362.700 Retinitis pigmentosa
      190.500 Retinoblastoma
      771.000 Rubella, congenital (in utero infections only)
    # 685.100 Sacral dimple
    # 216.910 Sebaceous cyst
# 282.600 Sickle cell anemia
Τ
    # 090.000 Syphilis, congenital (in utero infections only)
      238.030 Teratoma, abdomen
      238.010 Teratoma, head and face
      238.020 Teratoma, neck
      238.000 Teratoma, NOS
      238.080 Teratoma, other specified
      238.040 Teratoma, sacral, coccygeal
      257.800 Testicular feminization syndrome
      771.090 TORCH infection, unspecified (in utero infections only)
    # 608.200 Torsion of the testes or spermatic cord
      771.210 Toxoplasmosis (in utero infections only)
    # 553.100 Umbilical hernia
    # 286.400 von Willebrand disease
      335.000 Werdnig-Hoffman disease
      189.000 Wilms tumor (nephroblastoma)
      426.705 Wolfe-Parkinson-White syndrome, congenital
```

List ordered by 6-digit code number

```
# 090.000
                     Syphilis, congenital (in utero infections only)
                     Neoplasms of the liver
            155.000
                     Includes:
                                   hepatoblastoma
                                   hmangio-epithelioma
            159.800 Neoplasms of the abdomen
            162.800 Neoplasms of the lung
            171.800 Neoplasms of connective tissue
                     Includes:
                                   Ewing's sarcoma
                                   fibrosarcoma
            186.000 Neoplasms of the testes
            189.000 Wilms tumor (nephroblastoma)
            190.500 Retinoblastoma
            191.000 Neoplasms of the CNS
                     Includes:
                                   gliomas
                                   mdulloblastoma
            194.000 Neuroblastoma
            202.300 Histiocytosis, malignant
            208.000 Leukemia, congenital, NOS
      214
                     Lipoma
            214.000 Lipoma, skin and subcutaneous tissue of face
             214.100 Lipoma, other skin and subcutaneous tissue
            214.200 Lipoma, intrathoracic organs
            214.300 Lipoma, intra-abdominal organs
            214.400 Lipoma, spermatic cord
            214.800 Lipoma, other specified sites
            214.810 Lipoma, lumbar or sacral lipoma
                     paraspinal lipoma
            214.900 Lipoma, unspecified site
T
      216
                     Benign neoplasm of skin
                      (NOTE: All benign neoplasms should be coded ONLY if
                       another reportable code is present)
                                   blue nevus pigmented nevus
                      Includes:
                                   papilloma
                                               dermatofibroma
                                   syringoadenoma
                                   *dermoid cyst
                                   hydrocystoma
                                   syringoma
                     Excludes:
                                   skin of female genital organs (use 221.000),
                                   skin of male genital organs (use 222.000)
          # 216.000
                    Skin of lip
                     Excludes:
                                   vermillion border of lip
          # 216.100 Eyelid, including canthus
                                   cartilage of eyelid
                     Excludes:
          # 216.200 Ear and external auditory canal
                     Includes:
                                   auricle ear
                                   external meatus
                                   auricular canal
                                   external canal
                                   pinna
                                   cartilage of ear
                     Excludes:
```

S = Rev. 8/07 R = Rev. 6/07 N = Rev. 5/07 T = Rev. 6/04 * = code created b

```
216.400
                     Scalp and skin of neck
            216.500 Skin of trunk, except scrotum
                           Includes:
                                         axillary fold
                                         perianal skin
                              skin of:
                                         chest wall
                                         abdominal wall
                                         groin
                                         buttock
                                         anus
                                         perineum
                                         back
                                         umbilicus
                                         breast
                      Excludes:
                                   anal canal
                                   anus, NOS
                                   skin of scrotum
            216.600
                     Skin of upper limb, shoulder
            216.700
                     Skin of lower limb, hip
            216.800 Other specified sites of skin
                      Excludes:
                                   epibulbar dermoid cyst (use 743.810)
            216.900 Site unspecified
            216.910
                     Sebaceous cyst
             216.920 Hairy nevus
            221.000 Benign skin neoplasm of female genital organs
            222.000 Benign skin neoplasm of male genital organs
T
      228.0
                      Hemangioma
                      Include if greater than 4-inches diameter, if multiple
                      hemangiomas, or if cavernous hemangioma
            228.000
                      Hemangioma, of unspecified site
                      Always code if multiple hemangiomas of any size are present, if one
                      or more cavernous hemangiomas of any size are present, or if a single
                      hemangioma measuring ≥ 4cm in diameter or desribed as large, huge, or
                      of medical significance is present.
                     Hemangioma, skin & subcutaneous, NOS
            228.010
                      Always code if multiple hemangiomas of any size are present, if one
                      or more cavernous hemangiomas of any size are present, or if a single
                      hemangioma measuring ≥ 4cm in diameter or desribed as large, huge, or
                      of medical significance is present.
             228.020 Hemangioma, intracranial (Always code regardless of size, type or
                      number)
             228.030
                     Hemangioma, retinal (Always code regardless of size, type or number)
             228.040 Hemangioma, intra-abdominal (Always code regardless of size, type or
                      number)
             228.090 Hemangioma, of other sites (Always code regardless of size, type or
                      number)
             228.100 Cystic hygroma
                      Lymphangioma, any site
             237.700 Neurofibromatosis
             238.000 Teratoma, NOS
             238.010 Teratoma, head and face
             238.020 Teratoma, neck
             238.030 Teratoma, abdomen
             238.040 Teratoma, sacral, coccygeal
```

S = Rev. 8/07 R = Rev. 6/07 N = Rev. 5/07 T = Rev. 6/04 * = code created

```
# 243.990 Hypothyroidism, congenital
           (Exclude even if other defects are present only if the record
            specifies hypothyroidism of prematurity <36 weeks. Include other
            types of hypothyroidism and hypothyroidism NOS only when another
            reportable defect is present regardless of gestational age)
 251.200 Hypoglycemia, idiopathic
# 252.100 Hypoparathyroidism, congenital
  253.280 Hypopituitarism, congenital
  253.820 Diencephalic syndrome
  255.200 Adrenogenital syndrome (adrenal hyperplasia)
  257.800 Testicular feminization syndrome
 270.100 Phenylketonuria (PKU)
 270.200 Albinism
 270.300 Maple syrup urine disease
 270.600 Arginosuccinic aciduria
# 270.700 Hyperglycinemia
# 271.000 Glycogen storage diseases
# 275.330 Hypophosphatemic rickets
 277.000 Cystic fibrosis with no mention of meconium ileus
 277.010 Cystic fibrosis with mention of meconium ileus
  277.400 Disorders of bilirubin excretion
  277.510 Hurler syndrome
           Includes: lipochondrodystrophy
 277.620 Alpha-1 antitrypsin deficiency
 277.630 Pseudocholinesterase enzyme deficiency
  279.110 DiGeorge syndrome
  279.200 Combined immunodeficiency syndrome
 282.000 Hereditary spherocytosis
 282.100 Hereditary elliptocytosis
 282.200 Glucose-6-phosphate dehydrogenase (G-6-PD) deficiency
 282.600 Sickle cell anemia
 284.000 Red cell aplasia
 286.000 Hemophilia (all types)
 286.400 von Willebrand disease
 330.100 Cerebral lipidoses
          Includes:
                       Tay-Sachs disease
                        gangliosidosis
  331.890 Familial degenerative CNS disease
  335.000 Werdnig-Hoffman disease
  345.600 Infantile spasms, congenital
 351.000 Facial palsy
  352.600 Moebius syndrome
  362.600 Retinal degeneration, peripheral
  362.700 Retinitis pigmentosa
  363.200 Chorioretinitis
 368.000 Esotropia
 378.000 Exotropia
 379.500 Nystagmus
  425.300 Endocardial fibroelastosis
  426.705
          Congenital Wolfe-Parkinson-White syndrome
  427.900 Cardiac arrhythmias, NEC. Never code premature atrial
           contractions, PACs.
  453.000 Budd-Chiari, occlusion of hepatic vein
# 457.800 Other specified disorders of lymphatics (including chylothorax)
```

```
# 520.600 Natal teeth
      524.000 Abnormalities of jaw size
               micrognathia
               macrognathia
    * 524.080 Pierre Robin sequence
    # 550.000- Inguinal hernia or patent processus vaginalis never
       550.900 code in infants if <36 weeks gestation regardless of the
               presence of a reportable defect.
               NOTE: for those ≥36 weeks:
                Code in males only if another reportable defect is present;
                in females, always code even if found in isolation
     553.100 Umbilical hernia
      553.200 Epigastric hernia
    # 608.200 Torsion of testes or spermatic cord
   # 658.800 Amniotic bands (constricting bands, amniotic cyst)
    \# 685.100 Pilonidal sinus (sacrodermal), sacral sinus, sacral dimple
      760.710 Fetal alcohol syndrome
      760.718 Probable fetal alcohol syndrome
               Includes:
                            "facies"
      760.750 Fetal hydantoin (Dilantin) syndrome
    # 767.600 Erb's palsy
771
               Congenital infections (in utero infections only)
               Excludes: congenital syphilis (use 090.000)
      771.000 Rubella, congenital
      771.090 TORCH infection, unspecified
      771.100 Cytomegalovirus (CMV)
      771.210 Toxoplasmosis
      771.220 Herpes simplex
               Includes:
                           encephalitis
                            meningoencephalitis
      771.280 Congenital infection, other specified
                           human immunodeficiency virus (HIV) infection and
               Excludes:
                            acquired immunodeficiency syndrome (AIDS)
      774.480 Hepatitis, neonatal, other specified
      774.490 Hepatitis, neonatal, NOS
     777.100 Meconium plug syndrome
     777.600 Meconium peritonitis
     778.000 Ascites, congenital
    # 778.600 Hydrocele, congenital
```

HHS:PHS:CDC:NCBDDD:DBDDD:06/16/04 Doc. 6digit88, Version 06/04

EXCLUSION LIST for the MACDP Nonreportable birth defects

Conditions Never to be Reported

The following newborn and infant conditions include those descriptions considered to be excludable or nonreportable conditions in the MACDP. This includes certain biochemical disorders not considered part of the present MACDP case definition.

Alphabetical list of conditions that are never considered to be defects.

Description

Anal fissure Atrial contractions, premature Breast hypertrophy Bronchopulmonary dysplasia (Wilson-Mikity syndrome) Cephalohematoma Chalasia (gastroesophageal reflux) CNS hemorrhage Conjunctivitis Diastasis recti Epulis Gastroesophageal reflux Gum cysts - Includes epulis, ranula, mucocele Hydrocephalus secondary to intraventricular hemorrhage (IVH) or CNS bleed Hip click, with no follow-up or therapy T Heart murmur Hyaline membrane disease Intestinal obstruction - requires chart review to determine if T cause of obstruction is a reportable defect. If so, code only T Intussusception - requires chart review to determine if cause of intussusception is a reportable defect. If so, code only the cause. Inverted nipples Laryngotracheomalacia or tracheomalacia Meconium stained skin or nails Mucocele Neonatal acne Overriding (overlapping) sutures Petechiae Phimosis Pneumothorax Premature atrial contractions Protruding tongue Ranula Redundant foreskin Retractile testes Tracheomalacia T Volvulus - requires chart review to determine if cause of volvulus is a reportable defect. If so, code only the cause. Wilson-Mikity syndrome R = Rev. 6/07

N = Rev. 5/07T = Rev. 6/04* = code created by CDC # = on the MACDP Excl List

EXCLUSION LIST for the MACDP Nonreportable birth defects

Conditions Which may be Included Under Certain Conditions

The following newborn and infant conditions include those descriptions considered to be excludable or nonreportable conditions in the MACDP, <u>but which may be included</u> under certain circumstances.

The following rules apply to coding these conditions:

- A. If a condition or defect listed appears in a chart, singly or in any combination with other defects listed only on the Exclusion List, <u>do not</u> fill out the case record form.
- B. If one of these conditions listed accompanies a reportable birth defect (from the 6-digit code manual and not on the exclusion list), then use the listed 6-digit code and record <u>all</u> defects (including those from this list) from the hospital record onto the case abstraction form.

<u>Alphabetical</u> list of conditions requiring \underline{no} record abstraction to be performed \underline{unless} associated with a reportable defect. The addition or revision dates of the changes in the list of conditions requiring no record abstraction are shown.

| Revised/ |
|----------|
| Changed |
| Date |

| Changed Date | | | Code | Description |
|-----------------|-------|---|--|---|
| | | | 744.100 757.650 270.200 277.620 | Accessory auricle Accessory nipple (supernumerary nipple, or skin tag) Albinism Alpha 1-antitrypsin deficiency |
| | T | | 658.800 757.310 | Amniotic bands (constricting bands, amniotic cyst) Anal tags |
| 10/1/92 | T | | 746.400 270.600 | Aortic valve insufficiency or regurgitation, congenital - Code cases designated as 'mild', minimal', 'trivial', or 'physiologic' only if another reportable defect is present. Code all other degrees of insufficiency or regurgitation, including those where the degree is not specified, regardless of whether another reportable defect is present. Argininosuccinic aciduria |
| | T | | 778.000 744.220 | Ascites or anasarca, congenital. Includes: hydrops fetalis Bat ear |
| | T | # | 216.200 | Benign neoplasm of skin, ear and external auditory canal Includes: auricle ear external meatus auricular canal external canal pinna Excludes: cartilage of ear |
| | T | # | 216.100 | Benign neoplasm of skin, eyelid, including canthus |
| | | | | Excludes: cartilage of eyelid |
| | T | # | 216.000 | Benign neoplasm of skin, lip Excludes: vermillion border of lip |
| | D C/O | 7 | | |

^{* =} code created by CDC # = on the MACDP Excl List

EXCLUSION LIST for the MACDP Nonreportable birth defects

Alphabetical - Conditions Which may be Included Under Certain Conditions

| Revised/ Changed | , | | | |
|---------------------|---|---|--------------------|---|
| Date | | | Code | Description |
| | T | # | 216.700 | Benign neoplasm of skin, lower limb, hip |
| | T | # | 216.300 | Benign neoplasm of skin, other and unspecified parts of face Includes: cheek, external nose, external eyebrow, temple |
| | T | # | 216.800 | Benign neoplasm of skin, other specified sites of skin Excludes: epibulbar dermoid cyst (use 743.810) |
| | T | # | 216.400 | Benign neoplasm of skin, scalp and skin of neck |
| | T | # | 216.900 | Benign neoplasm of skin, site unspecified |
| | | | 216.500 | Benign neoplasm of skin, trunk, except scrotum Includes: axillary fold perianal skin skin of: chest wall, abdominal wall, groin, buttock, anus, |
| | | | | perineum, back, umbilicus, breast |
| | | | | Excludes: anal canal, anus, NOS skin of scrotum |
| | | # | 216.600 | Benign neoplasm of skin, upper limb, shoulder |
| | | | 221.000 | Benign skin neoplasm of female genital organs |
| | | | 222.000 | Benign skin neoplasm of male genital organs |
| | | | 754.020 | Bent nose, deviation of nasal septum |
| | | | 744.820 | Big lips Birth mark, NOS |
| | | | 757.385 743.450 | Blue sclera - if <36 weeks gestation, code only if |
| | | | 743.430 | another reportable defect is present. Always code if |
| | | | | ≥36 weeks gestation. |
| | | | 743.800 | Brushfield spots |
| | | | 757.390 | Cafe au lait spots |
| | | | 746.860 | Cardiomegaly, congenital NOS |
| | | | 744.230 | Cauliflower ear |
| | | | 330.100 | Cerebral lipidoses (e.g., Tay-Sachs, gangliosidoses, etc.) |
| | | | 756.200 | Cervical rib |
| | | | 755.500 | Clinodactyly (incurving of fifth finger) |
| 1/1/93 | | | 752.520 | Cryptorchidism (see undescended testicle) |
| | | | 277.010 | Cystic fibrosis, with mention of meconium ileus |
| | | | 277.000 | Cystic fibrosis, with no mention of meconium ileus |
| | | | 744.280 | Darwin's tubercle |
| 1/1/96 | T | | 754.030 | Dolichocephaly - if <36 weeks gestation, code only if another reportable defect is present. Always code if ≥36 weeks gestation. |
| 1/1/93 | | | 743.800 | Downward eye slant (antimongoloid) |
| | | | 744.110 | Ear tags, preauricular |
| | | | 744.120 | Ear tags, other |
| | | | 744.230 | Elfin ear, absent or decreased ear cartilage - if <36 weeks gestation, code only if another reportable defect is present. |
| | | | 743.800 | Epicanthal folds |
| | | | 767.600 | Erb's palsy |

EXCLUSION LIST for the MACDP Nonreportable birth defects

Erb's palsy

R = Rev. 6/07 N = Rev. 5/07 T = Rev. 6/04 * = code created by 767.600

T = Rev. 6/04

* = code created by CDC

= on the MACDP Excl List

Revised/

Alphabetical - Conditions Which may be Included Under Certain Conditions

| Changed | | | |
|---------|---|---------|---|
| Date | | Code | Description |
| | | 368.000 | Esotropia |
| | | 378.000 | Exotropia |
| | | 351.000 | Facial palsy |
| | | 757.380 | Flammeus nevus or port wine stain |
| | | 748.180 | Flat bridge of nose |
| | | 754.040 | Fontanelle (large or small) |
| | T | 743.630 | Fused eyelids - never code if <25 weeks gestation unless another reportable defect is present |
| | | 752.440 | Fusion of vulva |
| | | 282.200 | Glucose-6-phosphate dehydrogenase (G-6-PD) deficiency |
| | | 271.000 | Glycogen storage disease |
| | | 746.990 | Heart murmur - if chart review does not confirm a |
| | | | heart defect within 6 months, do not code as a defect even if other codable defects are present |
| | | 286.000 | Hemophilia |
| | | 751.620 | Hepatomegaly |
| | | 282.100 | Hereditary elliptocytosis |
| | | 282.000 | Hereditary spherocytosis |
| 3/4/91 | | 750.240 | High arched palate |
| | | 778.600 | Hydrocele, congenital |
| | | 752.480 | Hymenal tags |
| | | 270.700 | Hyperglycinemia |
| | | 251.200 | Hypoglycemia, idiopathic |
| | | 252.100 | Hypoparathyroidism, congenital |
| | | 275.330 | Hypophosphatemic rickets |
| 1/1/96 | T | 752.440 | Hypoplastic labia majora - if <36 weeks gestation, code only if another reportable defect is present. Always code if ≥ 36 weeks gestation. |
| 3/4/91 | T | 748.510 | Hypoplasia of lung; pulmonary hypoplasia - exclude only if an isolated defect in infants <36 weeks gestation |
| | | 752.810 | Hypoplastic scrotum - exclude if secondary to undescended testes |
| | T | 243.990 | Hypothyroidism, congenital (Exclude hypothyroidism of prematurity in infants <36 weeks gestation even if |

550.000Inguinal hernia or patent processus vaginalis. Never
code in infants <36 weeks gestation regardless of the
presence of a reportable defect. For infants ≥36
weeks:
In males, code only if another reportable defect is
present;
In females, always code even if found in isolation
Lanugo, excessive or persistent
Large fontanelle

other reportable defects are present. Include other types of hypothyroidism and hypothyroidism NOS when another reportable defect is present regardless of

EXCLUSION LIST for the MACDP Nonreportable birth defects

gestational age)

Imperforate hymen

Incurving fingers (clinodactyly)

R = Rev. 6/07 N = Rev. 5/07 T = Rev. 6/04 * = code created by CDC # = on the MACDP Excl List

Τ

752.430

755.500

Alphabetical - Conditions Which may be Included Under Certain Conditions

| Revised/ Changed | | | |
|---------------------|-----|--------------------|--|
| Date | | Code | Description |
| | | 755.500 | Long fingers and toes |
| | | 744.230 | Lop ear |
| | | 744.245 | Low set ears |
| | | 744.820 | Macrocheilia (big lips) |
| | | 270.300 751.010 | Maple syrup urine disease Meckel's diverticulum |
| | | 777.600 | Meconium peritonitis |
| | | 777.100 | Meconium plug |
| 9/10/90 | | 754.520 | Metatarsus varus or adductus |
| 10/1/00 | T | 744.830 | Microcheilia (small lips) |
| 10/1/92 | T | 746.600 | Mitral valve insufficiency or regurgitation, congenital - |
| | | | Code cases designated as 'mild', minimal', 'trivial', |
| | | | or 'physiologic' only if another reportable defect is |
| | | | present. Code all other degrees of insufficiency or |
| | | | regurgitation, including those where the degree is not |
| | | | specified, regardless of whether another reportable defect is present. |
| | | 757.386 | Mongolian spots |
| | | 743.650 | Nasal lacrimal duct obstruction |
| | | 520.600 | Natal teeth |
| | | 745.500 | Nonclosure of foramen ovale, NOS (see PFO) |
| 0 /4 0 / 0 0 | | 379.500 | Nystagmus |
| 9/10/90 | | 756.080 | Occiput, flat or prominent |
| 3/5/90 | | 457.800 | Other specified disorder of lymphatics, including |
| | | 755.600 | chylothorax Overlapping toes |
| 10/14/92 | Т | 747.000 | Patent ductus arteriosus (PDA) |
| 10/14/32 | 1 | 747.000 | 1)Always code if \geq 36 weeks of gestation and defect last |
| | | | noted at ≥ 6 weeks of age. 2)If ≥ 36 weeks gestation and defect last noted <6 weeks |
| | | | of age, code only if the PDA was treated)e.g. by |
| | | | ligation or indomethicin) or if another reportable |
| | | | defect is present. |
| | | | 3) Never code if <36 weeks gestation or if treated with |
| / / | | | prostaglandins regardless of gestational age. |
| 10/14/92 | T # | 745.500 | Nonclosure of foramen ovale, NOS |
| | | | Patent foramen ovale (PFO) 1)Always code if ≥36 weeks of gestation and defect last |
| | | | noted at ≥ 6 weeks of age. |
| | | | 2) If ≥36 weeks gestation and defect last noted <6 weeks |
| | | | of age, code only if another reportable defect is |
| | | | present. |
| | | | 3) Never code if <36 weeks gestation regardless of presence of other defects. |
| | T | 753.700 | Patent urachus |
| 0 /1 /00 | | 744.820 | Patulous lips (wide lips) |
| 8/1/93 | | 747.325 | Peripheral pulmonic stenosis (PPS) murmur - \underline{do} collect if PPS documented by echocardiogram |
| | | 270.100 | Phenylketonuria (PKU) |
| | | | EXCLUSION LIST for the MACDP |

^{* =} code created by CDC # = on the MACDP Excl List

Nonreportable birth defects

Alphabetical - Conditions Which may be Included Under Certain Conditions

| Revised/ Changed | | | |
|---------------------|--------|---|--|
| <u>Date</u> | | Code 685.100 744.230 744.230 755.006 | Description Pilonidal or sacral dimple Pixie-like ear Pointed ear Polydactyly in blacks (postaxial, type B), includes only skin tags on hands or feet. All other types of postaxial polydactyly (i.e. extra finger with bone, nail, etc.) should always be coded. Posteriorly rotated ears |
| | T | 744.410 744.110 747.680 752.450 | Preauricular sinus, cyst or pit Preauricular tags Primary pulmonary artery hypertension Prominent clitoris |
| | | 277.630 | Pseudocholinesterase enzyme deficiency |
| 10/1/92 | T | 750.500 751.580 284.000 744.500 755.616 685.100 | Pulmonary valve insufficiency or regurgitation, congenital Code cases designated as 'mild', minimal', 'trivial', or 'physiologic' only if another reportable defect is present. Code all other degrees of insufficiency or regurgitation, including those where the degree is not specified, regardless of whether another reportable defect is present. Pylorospasm (intermittent pyloric stenosis) Rectal fissures Red cell aplasia Redundant neck skin folds Rocker-bottom feet Sacral dimple |
| 1/1/96 | T # | 754.060 | Scaphocephaly, no mention of craniosynostosis Always code if ≥36 weeks gestation. If <36 weeks gestation, code only if another reportable |
| | | | defect is present. |
| 1/1/96 | T | 216.910 744.900 282.600 757.200 757.200 747.500 757.390 754.040 744.830 | Sebaceous cysts Short neck Sickle cell anemia Sidney line Simian crease (transverse palmar crease) Single umbilical artery Skin cysts Small fontanelle Small lips |
| 1/1/96 | T # | 757.640 | Small nipple (hypoplastic) Always code if ≥36 weeks gestation. If <36 weeks gestation, code only if another reportable defect is present |
| 10/1/92 7/13/92 | T | 759.020 090.000 759.240 755.630 750.000 | Splenomegaly Syphilis, congenital Thymic hypertrophy Tibial torsion Tongue-tie EXCLUSION LIST for the MACDP |

^{* =} code created by CDC # = on the MACDP Excl List

Nonreportable birth defects

Alphabetical - Conditions Which may be Included Under Certain Conditions

| Revised/ Changed | | | |
|---------------------|---|--|--|
| Date | | Code | Description |
| | | 608.200 | Torsion of spermatic cord |
| | _ | 608.200 | Torsion of testes |
| 10/1/92 | T | 746.105 | Tricuspid valve insufficiency or regurgitation, congenital - |
| | | | Code cases designated as 'mild', minimal', 'trivial', or 'physiologic' only if another reportable defect is present. Code all other degrees of insufficiency or regurgitation, including those where the degree is not specified, regardless of whether another reportable defect is present. |
| | | 759.900 | Umbilical cord atrophy |
| | | 553.100 | Umbilical hernias (completely covered by skin) |
| 1/1/93 | T | 752.500- | Undescended testicle (cryptorchidism) |
| | Т | 752.520 748.180 | 1) If < 36 weeks gestation, code only if there is a medical/surgical intervention for this problem; 2) If ≥36 weeks gestation and defect last noted at <1 year of age, code only if there was a medical/surgical intervention for this problem or if another reportable defect is present 3) Always code if ≥36 weeks gestation and defect first noted at ≥1 of age. |
| | | 748.180 | Upturned nose Upward eye slant (mongoloid) |
| | | 752.460 | Vaginal cysts |
| | | 752.480 | Vaginal tags |
| - 4 4 | _ | 286.400 | von Willebrand's disease |
| 3/14/91 | T | 755.130 | Webbed toes Code webbing of the second and third toes only if another reportable defect is present. Always code webbing of other toes regardless of whether another reportable defect is present |
| | | 744.500 748.180 755.600 757.680 | Webbing of neck Wide nasal bridge Widely spaced first and second toes Widely spaced nipples |

EXCLUSION LIST for the MACDP

 $\underline{\underline{\textbf{Numerical}}}$ list of conditions requiring \underline{no} record abstraction $\underline{\underline{\textbf{unless}}}$ associated with a reportable defect. The addition or revision dates of the changes in the list of conditions requiring no record abstraction are shown. .

Revised/ Changed

| Date Date | | Code | Description |
|-----------|-------------|-------------------------------|--|
| 7/13/92 | 216 | 090.000 | Syphilis congenital Benign neoplasm of skin (NOTE: All benign neoplasms should be coded ONLY if another reportable code is present) Includes: blue nevus pigmented nevus papilloma dermatofibroma syringoadenoma *dermoid cyst hydrocystoma |
| | | Exclude | syringoma s: skin of female genital organs (use 221.000), skin of male genital organs (use 222.000) |
| | # | 216.000 | Skin of lip |
| | # | 216.100 | Excludes: vermillion border of lip Eyelid, including canthus Excludes: cartilage of eyelid |
| | # | 216.200 | Ear and external auditory canal Includes: auricle ear |
| | # | 216.300 | Excludes: cartilage of ear Skin of other and unspecified parts of face |
| | | | <pre>Includes: cheek,external nose, external eyebrow, temple</pre> |
| | # | 216.400 | Scalp and skin of neck |
| | # | 216.500 | Skin of trunk, except scrotum Includes: axillary fold |
| | # # # | 216.600 216.700 216.800 | Excludes: anal canal anus, NOS skin of scrotum Skin of upper limb, shoulder Skin of lower limb, hip Other specified sites of skin Excludes: epibulbar dermoid cyst (use 743.810) |

^{* =} code created by CDC # = on the MACDP Excl List

216.900 Site unspecified **EXCLUSION LIST** for the MACDP

 $\frac{\textbf{Numerical}}{\textbf{a reportable defect.}} \text{ list of conditions requiring } \underline{\textbf{no}} \text{ record abstraction } \underline{\textbf{unless}} \text{ associated with a reportable defect.} \text{ The addition or revision dates of the changes in the list of conditions requiring no record abstraction are shown.}.$

Revised/ Changed

| Changed | | | |
|-------------|---------|----------|---|
| <u>Date</u> | | Code | <u>Description</u> |
| | | | |
| | # | 216.910 | Sebaceous cyst |
| | | 221.000 | Benign skin neoplasm of female genital organs |
| | | 222.000 | Benign skin neoplasm of male genital organs |
| | T | 243.990 | Hypothyroidism, congenital (Exclude even if other |
| | | | defects are present only if the record specifies |
| | | | hypothyroidism of prematurity <36 weeks. Include other |
| | | | types of hypothyroidism and hypothyroidism NOS only |
| | | | when another reportable defect is present regardless |
| | | | of gestational age. |
| | | 251.200 | Hypoglycemia, idiopathic |
| | | 252.100 | Hypoparathyroidism, congenital |
| | | 270.100 | Phenylketonuria (PKU) |
| | | 270.200 | Albinism |
| | | 270.300 | Maple syrup urine disease |
| | | 270.600 | Argininosuccinic aciduria |
| | | 270.700 | Hyperglycinemia |
| | | 271.000 | Glycogen storage diseases |
| | | 275.330 | Hypophosphatemic rickets |
| | | 277.000 | Cystic fibrosis, with no mention of meconium ileus |
| | | 277.010 | Cystic fibrosis, with mention of meconium ileus |
| | | 277.620 | Alpha 1-antitrypsin deficiency |
| | | 277.630 | Pseudocholinesterase enzyme deficiency |
| | | 282.000 | Hereditary spherocytosis |
| | | 282.100 | Hereditary elliptocytosis |
| | | 282.200 | Glucose-6-phosphate dehydrogenase (G-6-PD) deficiency |
| | | 282.600 | Sickle cell anemia |
| | | 284.000 | Red cell aplasia |
| | | 286.000 | Hemophilia |
| | | 286.400 | von Willebrand's disease |
| | | 330.100 | Cerebral lipidoses (e.g., Tay-Sachs, gangliosidoses, |
| | | 251 000 | etc.) |
| | | 351.000 | Facial palsy |
| | | 368.000 | Esotropia |
| | | 378.000 | Exotropia |
| 2/5/00 | | 379.500 | Nystagmus |
| 3/5/90 | | 457.800 | Other specified disorder of lymphatics, including |
| | | 520.600 | chylothorax Natal teeth |
| | т | | |
| | T | 550.000- | Inguinal hernia or patent processus vaginalis never |
| | | 550.900 | code in infants if <36 weeks gestation regardless of |
| | | | the presence of a reportable defect. NOTE: for those ≥36 weeks: |
| | | | |
| | | | in males, code only if another reportable defect is |
| | | | present; |
| | | EE2 100 | in females , always code even if found in isolation |
| | | 553.100 | Umbilical hernias (completely covered by skin) |
| R = Re | v 6/07 | 608.200 | Torsion of spermatic cord |
| | v. 6/07 | | |

^{* =} code created by CDC # = on the MACDP Excl List

Revised/

608.200 Torsion of testes

EXCLUSION LIST for the MACDP Nonreportable birth defects

Numerical - Conditions Which may be Included Under Certain Conditions

| Revised/ | | | |
|----------|---|------------|---|
| Changed | | <i>-</i> 1 | |
| Date | | Code | <u>Description</u> |
| | T | 658.800 | Amniotic bands (constricting bands, amniotic cyst) |
| | | 685.100 | Pilonidal or sacral dimple |
| | T | 743.450 | Blue sclera - if <36 weeks gestation, code only if |
| | | | another reportable defect is present. Always code if |
| | | | ≥36 weeks gestation. |
| | T | 743.630 | Fused eyelids - never code if <25 weeks gestation |
| | | | unless another reportable defect is present |
| | | 743.650 | Nasal lacrimal duct obstruction |
| | | 743.800 | Brushfield spots |
| | | 743.800 | Downward eye slant (antimongoloid) |
| | | 743.800 | Epicanthal folds |
| | | 743.800 | Upward eye slant (mongoloid) |
| | | 744.100 | Accessory auricle |
| | | 744.110 | Ear tags, preauricular |
| | | 744.120 | Ear tags, other |
| | | 744.220 | Bat ear |
| | | 744.230 | Cauliflower ear |
| | | 744.230 | Elfin ear, absent or decreased ear cartilage |
| | | | If <36 weeks gestation, code only if another |
| | | | reportable defect is present. |
| | | 744.230 | Lop ear |
| | | 744.230 | Pixie-like ear |
| | | 744.230 | Pointed ear |
| | | 744.245 | Low set ears |
| | | 744.246 | Posteriorly rotated ears |
| | | 744.280 | Darwin's tubercle |
| | | 744.410 | Preauricular sinus, cyst or pit |
| | | 744.500 | Redundant neck skin folds |
| | | 744.500 | Webbing of neck |
| | | 744.820 | Macrocheilia (big lips) |
| | | 744.820 | Patulous lips (wide lips) |
| | | 744.830 | Microcheilia (small lips) |
| | | 744.900 | Short neck |
| | | 745.500 | Nonclosure of foramen ovale, NOS (see PFO) |
| 10/14/92 | T | 745.500 | Patent foramen ovale (PFO) |
| | | | 1)Always code if ≥36 weeks of gestation and defect last |
| | | | noted at \geq 6 weeks of age. |
| | | | 2)If ≥36 weeks gestation and defect last noted <6 weeks |
| | | | of age, code only if another reportable defect is |
| | | | present. |
| | | | 3)Never code if <36 weeks gestation regardless of |
| | | | presence of other defects. |
| 10/1/92 | T | 746.020 | Pulmonary valve insufficiency or regurgitation, |
| | | | congenital - |
| | | | Code cases designated as 'mild', minimal', 'trivial', |
| | | | or 'physiologic' only if another reportable defect is |
| | | | present. Code all other degrees of insufficiency or |
| | | | regurgitation, including those where the degree is not |
| | | | |

^{* =} code created by CDC # = on the MACDP Excl List

specified, regardless of whether another reportable defect is present. $\,$

EXCLUSION LIST for the MACDP Nonreportable birth defects

Numerical - Conditions Which may be Included Under Certain Conditions

| Revised/ Changed | | g . J . | |
|---------------------|---|---|--|
| <u>Date</u> | | Code | <u>Description</u> |
| 10/1/92 | T | 746.105 | Tricuspid valve insufficiency or regurgitation, congenital - Code cases designated as 'mild', minimal', 'trivial', or 'physiologic' only if another reportable defect is present. Code all other degrees of insufficiency or regurgitation, including those where the degree is not specified, regardless of whether another reportable defect is present. |
| 10/1/92 | T | 746.400 | Aortic valve insufficiency or regurgitation, congenital - Code cases designated as 'mild', minimal', 'trivial', or 'physiologic' only if another reportable defect is present. Code all other degrees of insufficiency or regurgitation, including those where the degree is not specified, regardless of whether another reportable defect is present. |
| 10/1/92 | T | 746.860 746.990 | Mitral valve insufficiency or regurgitation, congenital - Code cases designated as 'mild', minimal', 'trivial', or 'physiologic' only if another reportable defect is present. Code all other degrees of insufficiency or regurgitation, including those where the degree is not specified, regardless of whether another reportable defect is present. Cardiomegaly, congenital NOS Heart murmur - if chart review does not confirm a heart defect within 6 months, do not code as a defect even if other codable defects are present |
| 10/14/92 | T | 747.000 | Patent ductus arteriosus (PDA) 1) Always code if ≥36 weeks of gestation and defect last noted at ≥6 weeks of age. 2) If ≥36 weeks gestation and defect last noted <6 weeks of age, code only if the PDA was treated)e.g. by ligation or indomethicin) or if another reportable defect is present. 3) Never code if <36 weeks gestation or if treated with |
| 8/1/93 | | 747.325 747.500 747.680 778.000 748.180 | prostaglandins regardless of gestational age. Peripheral pulmonic stenosis (PPS) murmur - do collect if PPS documented by echocardiogram Single umbilical artery Primary pulmonary artery hypertension Ascites or anasarca. Includes: hydrops fetalis Flat bridge of nose |

^{* =} code created by CDC # = on the MACDP Excl List

748.180 Upturned nose 748.180 Wide nasal bridge

EXCLUSION LIST for the MACDP Nonreportable birth defects

Numerical - Conditions Which may be Included Under Certain Conditions

| Revised/ Changed | | | |
|---------------------|-----|---|--|
| | | Date | Code Description |
| 3/4/91 | T | 748.510 | Hypoplasia of lung; pulmonary hypoplasia - exclude if isolated defect in infants <36 weeks gestation. |
| 3/4/91 | | 750.000 750.240 750.500 751.010 751.580 751.620 752.430 | Tongue-tie High arched palate Pylorospasm (intermittent pyloric stenosis) Meckel's diverticulum Rectal fissures Hepatomegaly Imperforate hymen |
| | | 752.440 | Fusion of vulva |
| 1/1/96 | T | 752.440 | Hypoplastic labia majora - if <36 weeks gestation, code only if another reportable defect is present. Always code if ≥ 36 weeks gestation. |
| 3/4/91 | | 752.450 752.460 752.480 752.480 | Prominent clitoris Vaginal cysts Vaginal tags Hymenal tags |
| 1/1/93 | T | 752.500- | Undescended testicle (cryptorchidism) |
| 2, 2, 30 | T | 752.520 | 1) If < 36 weeks gestation, code only if there is a medical/surgical intervention for this problem; 2) If ≥36 weeks gestation and defect last noted at <1 year of age, code only if there was a medical/surgical intervention for this problem or if another reportable defect is present 3) Always code if ≥36 weeks gestation and defect first noted at ≥1 of age. |
| 1/1/93 | | 752.520 752.810 753.700 754.020 | Cryptorchidism (see undescended testicle) Hypoplastic scrotum - exclude if secondary to undescended testes Patent urachus Bent nose, deviation of nasal septum |
| 1/1/96 | T | 754.030 | Dolichocephaly - if <36 weeks gestation, code only if another reportable defect is present. Always code if ≥36 weeks gestation. |
| 1/1/93 1/1/96 | | 754.040 754.060 | Fontanelle (large or small) Scaphocephaly, no mention of craniosynostosis If <36 weeks gestation, code only if another reportable defect is present. |
| 1/1/93 | | 754.520 755.006 | Always code if ≥36 weeks gestation. Metatarsus varus or adductus Polydactyly in blacks (postaxial, type B), includes only skin tags on hands or feet. All other types of postaxial polydactyly (i.e. extra finger with bone, nail, etc.) should always be coded. |
| 3/14 | /91 | T 755.1 | |

another reportable defect is present. Always code webbing of other toes regardless of whether another reportable defect is present

EXCLUSION LIST for the MACDP Nonreportable birth defects

Numerical - Conditions Which may be Included Under Certain Conditions

| Revised/ |
|----------|
| Changed |

| | Date | | Code | Description |
|------|---------|---|---------|---|
| | | | 755.500 | Clinodactyly (incurving of fifth finger) |
| | | | 755.500 | Long fingers and toes |
| | | | 755.600 | Overlapping toes |
| | | | 755.600 | Widely spaced first and second toes |
| | | | 755.616 | Rocker-bottom feet |
| | | | 755.630 | Tibial torsion |
| | | | 756.080 | Occiput, flat or prominent |
| | | | 756.200 | Cervical rib |
| | | | 757.200 | Sidney line |
| | | | 757.200 | Simian crease (transverse palmar crease) |
| | | | 757.310 | Anal tags |
| | | | 757.380 | Flammeus nevus or port wine stain |
| | | | 757.385 | Birth mark, NOS |
| | | | 757.386 | Mongolian spots |
| | | | 757.390 | Cafe au lait spots |
| | | | 757.390 | Skin cysts |
| | | | 757.450 | Lanugo, excessive or persistent |
| | 1/1/96 | T | 757.640 | Small nipple (hypoplastic) |
| | | | | If <36 weeks gestation, code only if another |
| | | | | reportable defect is present. |
| | | | | Always code if ≥36 weeks gestation. |
| | 9/10/90 | | 757.650 | Accessory nipple (supernumerary nipple, or skin |
| tag) | | | | |
| | | | 757.680 | Widely spaced nipples |
| | | | 759.020 | Splenomegaly |
| | | | 759.240 | Thymic hypertrophy |
| | | | 759.900 | Umbilical cord atrophy |
| | | | 767.600 | Erb's palsy |
| | | | 777.100 | Meconium plug |
| | | | 777.600 | Meconium peritonitis |
| | | | 778.000 | Ascites or anasarca, congenital |
| | | | 778.600 | Hydrocele, congenital |

R = Rev. 6/07

N = Rev. 5/07

T = Rev. 6/04

^{* =} code created by CDC # = on the MACDP Excl List

MACDP Decision Tree for Determining Whether to Include Patent Ductus Arteriosus (PDA)

```
Is the child on
prostaglandins?
              ----> Yes ---->
                                           Never code
     ω
   No
     ω
What was the gestational
age of the child at birth? -----> < 36 wks ---> Never code
     ω
   \geq 36 wks
     ω
How old was the
child when defect
was last noted? ----> > 6 wks ---> Always code
    ω
   < 6 wks
     \omega
Has the PDA been
treated? (e.g., by
ligation or
                    ----> Yes ----> Always code
indomethicin)
     \omega
   No
Include only if another reportable heart defect is present.
```

= on the MACDP Excl List

R = Rev. 6/07 N = Rev. 5/07 T = Rev. 6/04 * = code created by CDC

MACDP Decision Tree for Determining Whether to Include Patent Foramen Ovale (PFO)

R = Rev. 6/07 N = Rev. 5/07 T = Rev. 6/04 * = code created by CDC # = on the MACDP Excl List

MACDP Decision Tree for Determining Whether to Include Peripheral Pulmonary Stenosis (PPS)

May 22, 1996