

Common Genetic ICD-9 Codes

General Prenatal Testing

V82.4	(Maternal)Postnatal scrn for chrn anomalies
V28.8	CVS- Other Specified Prenatal Test
V28.2	Other screening based on amniocentesis
V28.1	Amniocentesis for Elev. AFP
V28.0	Amniocentesis to r/o chromosome abnormality
V26.3	Preconceptional Genetic Counseling
V26.3	Genetic Counseling & Testing
796.5	Abnor finding on anten screening (MSAFP screening)
795.2	non-specific Abnormal karyotype
659.63	Elderly multigravida - AMA >G2
659.53	Elderly primigravida AMA G1

Chromosomal related

279.11	Di George syndrome
	Velo-cardio-facial syndrome
655.13	Fetal chromos abnormal- antepartum
655.14	Fetal chromos abnormal- postpartum
758.0	Down syndrome
758.1	Trisomy 13
758.2	Trisomy 18
758.3	Autosomal deletion synd
758.4	Balanced autosomal transloc
758.5	Autosomal anomaly nec
758.6	Turner Syndrome
758.7	Klinefelter's syndrome
758.89	other condition due to chromosome anomaly

Abnormal U/S findings:

228.1	cystic hygroma
348.0	choroid plexus cyst Cerebral Cysts
653.63	Hydrocephal fetus-antepart
653.64	Hydrocephal fetus-postpar
653.7 1	Sacrococcygeal teratoma
653.7	Other fetal abn causing disprop
653.70	Other fetal disproportion-unspe
655.0	Fetal cns malformation
655.8	Fetal abnormal nec
655.83	Fetal abnormality, antepartum complication
655.93	Fetal abnormal unspec- antepartum
655.94	Fetal abnormal nos-postpartum
656.3	Fetal distress
656.5	Poor fetal growth
656.6	Excessive fetal growth
656.7 1	Placental Abruption
656.7 2	Placental Anomalies
656.7	Other placental conditions
656.8	Fetal/placental problem nec
658.03	olioghydramnios, antepartum
658.1	Premature rupture membranes
740.0	Anencephaly
740.1	Craniorachischisis
741	Spina bifida
741.00	Spina bifida (unspec region) with hydro
741.01	spina bifida (cervical) with

	hydrocephalus
741.02	spina bifida (dorsal-thoracic) with hydrocephalus
741.03	spina bifida (lumbar) with hydrocephalus
741.9	Meningocele
741.90	spina bifida(unspec region); no hydrocephalus
741.91	spina bifida (cervical); no hydrocephalus
741.92	spina bifida (dorsal-thoracic);no hydrocephalus
741.93	spina bifida(lumbar);no hydrocephalus
742.0 1	Hydroencephalocele
742.0 2	Exencephaly
742.0	Encephalocele
742.1	Microcephaly
742.2 0	Holoprosencephaly
742.2 1	Hemimegalencephaly
742.2 2	Lissencephaly
742.2	reduct. deform. of brain
742.3 1	Dandy Walker Malformation
742.3 3	hydranencephaly
742.3	Congenital hydrocephalus
742.4	Other specified anomalies of brain
746.89	Congenital heart anomaly nec
749.10	cleft lip, unspecified
749.20	Cleft lip/palate, unspec
749.24	bilateral cleft lip and palate
756.79	Gastroschisis
756.79	Omphalocele
756.79	Oth congenital anomalies of abdomen wall
761.2	Oligohydramnios affecting NB/FETUS
789.30	Abdon/pelvic swell, mass or lump, unspefd site
789.5	ascites
793.6	Abnormal finding/ultrasound/ abdominal area
761.3	Polyhydramnios affect nb/fetus
746.9	unspec congenital heart anomaly no

Pregancy Loss / fetal demise

630	Hydatidiform mole
656.40	intrauterine death, unspecf
674.9	Unspecified Neonatal Death
631	Other abnormal product of conception
656.43	intrauterine death, antepartum IUFD
656.4	Intrauterine death
634.90	Spontaneous abortion
779.9	Stillbirth

Multiple miscarriages

646.30	Hab Abort, unspec
646.33	Habitual aborter-antepart
646.34	Habitual aborter-postpart
648.00	Diabetes in preg-unspec
648.1	Maternal diabetes

Multiple pregnancies

651.03	Twin pregnancy
651.1	Triplet pregnancy
651.4	Triplet pregnancy w/fetal loss
651.6	Other multiple preg w/fetal loss
651.2	Quadruplet pregnancy
651.30	twin preg w fetal loss (unspec)
651.5	Quadruplet preg w/fetal loss
659.4	Grand multiparity

Family/Genetic History

655.20	Hered.disease in fam poss. aff.fetus
V18.4	Family history, mental retardation
V19.7	Family history of consanguinity
V19.5	Family History of congenital anomalies
V19.8	Family History of other conditions
655.23	Fetus at risk for Herediarty/Genetic disorder

Other Reasons

655.41	susp.fetal dam.due to etoh
655.5	fetal damage d/t drug
655.53	Suspected Teratogen exposure
655.6	Radiation fetal damage
656.0	Fetal-maternal hemorrhage
656.10	RH Immu(unspec as to ep of care)
656.13	Rh isoimmun ante partum condition
656.23	Other isoimmun antepartum
658.4	Infection amniotic cavity
659.8	Indicat care lab/del nec
663	Umbilical cord complic
663.0	Prolapse of cord
663.1	Compress cord round neck
663.2	Cord entangle with compress nec
663.4	Short umbilical cord
663.6	Vascular lesions of cord
663.8	Umbilical cord complic nec
760.71	Maternal alcohol affect nb
760.71	Fetal Alcohol Syndrome
760.8	Maternal cond nec affect nb
760.9	Maternal cond nos affect nb
761	Maternal compl affect nb
761.0	Incompetent cervix affect nb
761.1	Premat rupt memb affect nb
762.0	Placenta previa affect nb
762.1	Placenta hem nec affect nb
762.2	Abnormal plac nec/nos affect nb
762.6	Umbil cond nec affect nb
762.7	Chorioamnionitis affect nb
764	Slow fetal growth/malnut
764.90-9	Fetal Growth Retardation, nos

Postnatal

MENTAL DISORDERS

299.00	Infantile/child Autism(current)
312.9	behavior disturbance
314.00	Attention Deficit Disorder
315.2	Delayed learning
315.3	Delayed speech
315.31	Devl language disor
315.39	speech delay
315.5	Mixed development disorder
315.8	Other delays in development
315.9	Developmental delay
317	mild mental retardaton
318.0	Moderate mental retardation
318.1	Severe mental retardation
318.2	Profound mental retardation
319	Mental retardation nos

NEUROLOGIC

343.9	Infantile Cerebral Palsy
345.9	Epilepsy, unspecified
358.8	benign congenital hypotonia

EYES/VISION

743	Congenital eye anomaly
743.0	Anophthalmos
743.06	Cryptophthalmos
743.10	Microphthalmos, unspcfd
743.30	Congenital cataract nos
743.45	Aniridia
743.61	Congenital ptosis, blepharophimosis,
743.63	Spec anomaly eyelid nec
743.69	Anomaly eyelid/orb nec
743.9	Eye anomaly nos

EARS / HEARING

389.11	sensory hearing loss
389.9	unspecified hearing loss
744.0	Ear anomaly with impair hear
744.01	Congenital absence ext ear
744.09	Ear anomaly nec/impair hear
744.29	Ear anomaly nec
744.3	Unspec anomaly of ear NOS

CARDIAC - CEREBROVASCULAR

425.3	Cardiomyopathy, congenital
429.3	Cardiomegaly
745.2	Tetralogy of fallot

ORAL

524.04	Micrognathia
524.06	Retrognathia
525.8	disorders of teeth /jaw

ABDOMINAL

553.1	Umbilical Hernia
553.20	ventral hernia, unspcfd
553.3	Diaphragmatic hernia
553.8	Hernia, specified site
789.1	hepatomegaly
789.5	ascites

GENITAL / FERTILITY/ REPRODUCTION

256.3	Other ovarian failure
257.2	cryptorchidism
257.8	androgen insensitivity
259.0	Delay sexual development
259.1	precocious sex.dev
606.0	Azoospermia
606.1	Oligospermia
606.9	Male Infertility, unspec
608.3	Atrophy of testis
608.89	male genital disorder
626.0	amenorrhea (primary)
628.9	female infertility, unspec
629.9	Habitual aborter, not preg
646.30	Hab Abort, unspec
646.33	Habitual aborter-antepart
646.34	Habitual aborter-postpart
752.51	Undescended testes
752.61	Hypospadias
752.7	Indeterminate sex
752.9	Genital organ anomal nos

SKIN

701.1	Ichthyosis, acquired
709.09	cafe au lait

SKULL/HEAD/BRAIN

740.0	Anencephaly
740.1	Craniorachischisis
741	Spina bifida
741.9	Meningocele
741.90	spina bifida(unspec region); no hydrocephalus
742.0	Encephalocele

742.01	Hydroencephalocele
742.1	Microcephaly
742.20	Holoprosencephaly
742.3	Congenital hydrocephalus
742.4	Other specified brain anomal
742.42	Lissencephaly
742.5	oth.spec.anom. spinal cord
742.59	Spinal cord anomaly nec
742.8	Nervous system anomal
742.9	Unsp anomaly-brain,spinal cord

FACE/NECK

749.00	cleft palate unspecified
749.03	Bilat cleft palate-compl
749.10	cleft lip, unspecified
749.11	Unilat cleft lip-compl
749.20	CL/palate, unspec
749.24	Bilat cleft lip and palate
754.0	Cong skull/face/jaw def
754.01	Micrognathia
745.5	Webbing of neck
744.89	Cong anomal face & neck
748.0	Choanal atreis
748.1	Nose anomaly nec
750.10	Tongue anomaly, nos
750.15	Congenital macroglossia
750.3	Cong esoph fistula/atresia
751.21	Imperforate anus
756.0	Anomaly skull/face bone
756.08	Macrocephaly
756.10	anomalies of spine, unspecf
756.17	Spina bifida occulta
756.19	Anomaly of spine nec

HEART

745	Cardiac septal clos anomaly
745.0	Common truncus
745.1	Transpos of great vessel
745.2	Tetralogy of fallot
745.4	Ventricular sept defect
746	Other congenital heart anomaly
746.0	Pulmonary valve anomaly
746.86	Congenital heart block
746.89	Congenital heart anomaly nec
747.0	Patent ductus arteriosus
747.10	Coarctation of aorta
747.20	Cong anomaly of aorta nos
747.22	Aortic atresia/stenosis
747.29	Congl anomaly of aorta nec
747.3	Pulmonary artery anomaly
747.64	hypertrophy of lower limb
747.8	Circulatory anomaly nec
747.81	Cerebrovasc anomaly
747.9	Circulatory anomaly nos

RENAL

753	Urinary system anomaly
753.0	Renal agenesis
753.12	polycystic kidney dis, nos
753.15	Renal dysplasia
753.29	Congenital hydronephrosis
753.3	Kidney anomaly nec
753.5	Bladder exstrophy
753.9	Urinary anomaly nos

LIMBS - SKELETAL

754.51	Clubbed Foot
754.51	Talipes equinovarus
754.69	Congenital valgus foot def nec
754.79	Congenital foot deform nec
754.81	Pectus excavatum
754.89	Fetal Akinesia Sequence
755	Other congen limb anomaly
755.00	Polydactyly, NOS
755.10	Syndactyly, multiple/nos
755.20	Reduc deform up limb nos
755.3	Reduct deform lower limb
755.4	Reduct deform limb nos
755.41	ectrodactyly
755.50	Upper limb anomaly nos
755.58	Congenital cleft hand
755.59	Upper limb anomaly nec
755.60	Lower limb anomaly nos
755.66	Anomaly of toes nec
755.67	Anomaly of foot nec
755.69	Lower limb anomaly nec
755.9	Congen limb anomaly nos
756.1	Macrocosmia
756.0	Anomaly skull/face bone
756.08	Macrocephaly
756.10	anomalies of spine, unspecf
756.17	Spina bifida occulta
756.19	Anomaly of spine nec

756.3	Rib sternum anomaly nec
756.4	Chondrodystrophy
756.41	Achondroplasia
756.43	Chondrodysplasia Punctata
756.44	Hypochondroplasia
756.45	Thanatophoric dysplasia
756.51	Osteogenesis imperfecta
756.6	Anomaly of diaphragm
756.79	Oth cong anom of abd wall
756.791	Gastroschisis
756.79	Omphalocele
757.1	I Ichthyosis
757.2	Dermatoglyphic anomaly
757.32	Cutis Marmorata Telangiectatica Congenita
757.31	Congen ectodermal dysplas
757.39	Other spec anom of skin
757.4	Hair anomaly nec
757.5	Nail anomaly nec

CHROMOSOMES

758.0	Down syndrome /+21
758.1	Trisomy 13 / +13
758.2	Trisomy 18 / +18
758.3	Autosomal deletion synd
758.4	Balanced autosomal transloc
758.5	Autosomal anomaly nec
758.6	Turner Syndrome / -X
758.7	Klinefelter's syndrome / XXY
758.81	Other cond.due to sex chrom
758.89	other chromosome anomaly condition
758.9	Oth chromosome anomaly unspecif
795.2	non-specific Abnormal karyotype
796.5	Abnor finding on anten screening

Congenital Anomaly / Syndromes

279.11	Di George syndrome VCF
284.0	Fanconi's Anemia
237.71	NF-1
237.72	NF-2
759.7	MCA - Multiple congenital anomaly
759.81	Prader-Willi syndrome
759.82	Marfan Syndrome
759.83	Fragile X syndrome
759.89	Congenital malformation.syn
759.9	Congenital anomaly nos

PHYSICAL GROWTH

243.0	congenital hypothyroidism
244.9	unspecified hypothyroidism
251.2	Hypoglycemia- unspecified
255.2	cong. adrenal hyperplasia
278.00	Obesity, unspecified
764.90-9	Fetal Growth Retardation, nos
765	Short gestat/low birthwt
779.82	cong hypotonia
779.9	Stillbirth
780.39	seizures/convulsions, NOS
781.0	Abn involuntary movements - dystonia
781.2	Ataxia (gait)
781.3	Lack of coordin-hypotonia
783.1	Abnormal Weight Gain
783.22	Underweight
783.4	Failure to thrive
783.4 1	short stature
783.4 2	Lack of Growth
783.40	lack of nml physiological dev, unspec
783.41	Failure to thrive
783.42	delayed milestones
783.43	Short Stature
783.9 1	Tall Stature
784.5	speech disorder, defect
786.03	Apnea

Familial / Genetic

V13.2	Personal Hx of genital or obsteric disorder
V13.5	Personal Hx other musculoskeletal
V13.69	personal Hx of other congenital malformations
V18.4	FAM Hx, mental retardation
V19.0	FAM Hx Blindness
V19.5	FAM Hx of congenital anomalies
V19.7	FAM Hx of consanguinity
V19.8	FAM Hx of other conditions
V26.3 1	Preconceptional Genetic Counseling
V40.0	Problems with learning
V71.8	observ. of specif/susp cond.
V71.89	Observ for other specified susp cond
V71.9	observ for unspecf suspec condition

Solid Tumors & Neoplasms

200.10	Lymphosarcoma unspec
155.0	Hepatoblastoma
171.0	Malignant neoplasm of head, face, and neck
171.5	Malignant neoplasm abdomen
171.8	Malignant neoplasm of other specified sites of connective and other soft tissue
171.9	Malignant neoplasm of connective and other soft tissue, site unspecified
173.9	Other malignant neoplasm of skin, site unspecified
183.0	Malig neoplasm Ovary
186.0	Malig neoplasm of undes. testis
186.9	malig neopl of oth & unspc testis
188.0	Malignant neoplasm of trigone of urinary bladder
188.8	Malignant neoplasm of other specified sites of bladder
188.9	Malignant neoplasm of bladder, Retinoblastoma
190.5	Retinoblastoma
198.81	Secondary malignant neoplasm of breast
198.89	Secondary malignant neoplasm of other specified sites
198.89	Neoplasm of connective tissue-leg,malignant
199.1	Malig neoplasm nos,unsp site
200.00	Reticulosarcoma unspec
202.81	Other lymphomas; lymph nodes of face, head and neck
202.82	Other lymphomas; intrathoracic lymph nodes
202.83	Other lymphomas; intra-abdominal lymph nodes
202.84	Other lymphomas; lymph nodes of axilla and upper limb
202.85	Other lymphomas; lymph nodes of inguinal region and lower limb
202.86	Other lymphomas; intrapelvic lymph nodes
202.88	Other lymphomas; lymph nodes of multiple sites
203.00	Multiple myeloma/immunoproliferative neoplasms; no remission
203.01	Multiple myeloma and immunoproliferative neoplasms; remission
223.3	Benign neoplasm of bladder
233.0	Carcinoma in situ of breast
233.7	Carcinoma in situ of bladder
236.2	neoplasm uncertain behavior ovary
236.4	neoplasm uncertain behavior testis
236.7	Neoplasm of uncertain behavior of bladder
239.2	Neoplasm of unspecified nature of bone, soft tissue, and skin
239.3	Neoplasm unspecified nature

	of breast
239.3	neoplasms of the breast
239.4	neoplasms of the bladder

Leukemias

200.20	Burkitt's tumor unspec sites
200.21	BL (nodes of head, face, neck)
200.28	BL mult sites
200.80	Mixed lymphosarcoma unspec
201.00	Hodgkins without remission
201.90	- disorder nos unspec
202.00	Nodular lymphoma unspec
202.10	mycosis fungoides
202.20	Sezary's disease, unspec site
202.3	Malignant histiocytosis
202.30	Malignant histiocytosis, unspec
202.40	Hairy-cell leukem unspec
202.60	malig mast cell tumors, unsp site
202.80	Lymphomas nec unspec site
202.9	Malignant neoplasm lym/hist tis nec
202.90	Lymphoid mal nec unspec
203.00	Multiple Myeloma (<i>no remission</i>)
203.01	Mult. myeloma (<i>remission</i>)
203.10	Plasma Cell leuk, (<i>not in remission</i>)
203.11	Plasma cell leuk, in remi
203.8	Immunoproliferat neoplasm nec
203.80	immunprolif neoplas, (<i>no remission</i>)
204.00	ALL (<i>not in remission</i>)
204.01	ALL (<i>in remission</i>)
204.10	CLL (<i>not in remission</i>)
204.11	CLL (<i>in remission</i>)
204.2	Subac lymphoid leukemia
204.80	Other lymph leukemia (<i>no remission</i>)
204.81	Other lymph leuk (<i>in remission</i>)
204.90	Uspec. lymph leuk(<i>not in remission</i>)
204.91	unspec lymph leuk (<i>in remission</i>)
205.00	AML/APL Myeloid (<i>no remission</i>)
205.01	AML (<i>in remission</i>) APL Myeloid
205.10	CML (<i>not in remission</i>)
205.11	CML (<i>in remission</i>)
205.2	Subacut myeloid leukemia
205.30	Myeloid sarcoma (<i>not in remission</i>)
205.80	Myeloid leukemia nec (<i>not in remission</i>)
205.82	Myeloid leukemia (<i>in remission</i>)
205.90	Myeloid leukemia nos (<i>noremision</i>)
205.91	unspec myel leuk (<i>in remission</i>)
206.00	Acute Leukemia
206.10	Chronic monocytic leukemia (<i>not in remission</i>)
206.20	Subac monocytic leukemia (<i>not in remission</i>)
206.80	Monocytic leukemia nec (<i>not in remission</i>)

206.90	Monocytic leukemia nos (<i>not in remission</i>)
207.00	Other specific Leukemia
207.10	Chronic erythremia (<i>not in remission</i>)
207.20	Megakaryocytic leukemia (<i>not in remission</i>)
207.80	other specific leukemia (<i>not in remission</i>)
207.81	other specific leukemia (<i>in remission</i>)
208.0	Acute leukemia nos
208.00	Acute Leukemia (<i>not in remission</i>)
208.01	Acute Leukemia (<i>in remission</i>)
208.10	Chronic leukemia nos (<i>not in remission</i>)
208.2	Subacute leukemia nos
208.80	other leukemia of unspecified type, (<i>not in remission</i>)
208.81	oth leuke unspec type, in rem
208.9	Leukemia-unspec cell nos
214.0	Lipoma
215.0	benign neoplasm connective tissue, head and neck
215.4	benign neoplasm conn tissue and soft tissue- thorax
215.9	benign neoplasm conn tiss or other soft tiss, unspec
228.00	Hemangioma
238.4	PCV Polycythemia vera
238.6	Myeloma (multiple)
238.7	MDS Myelodysplastic/ myeloproliferative syndrome
284.8	Acquired aplas anem/pancytopenia
284.9	Aplastic anemia, unspec idiopathic
285.0	sideroblastic anemia
285.22	Anemia in neoplastic disease
285.9	anemia, unspec
287.1	thrombocytopenia
287.5	Thrombocytopenia, unspecified
288.0	Leukopenia, neutropenia
288.9	Other specified disease of WBCs (Leukocytosis)
758.89	other condition due to chromosome anomaly
758.9	Oth chromosome anomly unpecific
789.1	Hepatomegaly
789.2	Splenomegaly
789.30	Abdon/pelvic swell, mass or lump, unspcfd site
790.09	other abnormality of red blood cells
V10.62	Myeloid leukemia
V10.63	Monocytic leukemia
V10.69	Other leukemia
V10.71	lymphosarcoma and reticulosarcoma
V10.72	Hodgkin's disease

