

Confidential Birth Defect Registry Report

Instructions: This report form is to be used by physicians, pediatric specialty clinics and hospitals to report birth defects for children up to age two. The information is submitted to the Wisconsin Department of Health Services, Bureau of Community Health Promotion, Birth Defect Prevention and Surveillance Program.

Complete as much information as possible. Leave items **blank** if you do not have the information. Do not write "N/A" or similar in the spaces.

This report can be submitted via a secure website. Email dhswbdr@dhs.wisconsin.gov to request access to the site. If completing the report on paper, fax to Wisconsin Birth Defect Registry (WBDR), State administrator, Bureau of Community Health Promotion at 608-267-9042.

Parent or guardian consent is not required for reporting. Removal of names and addresses of the child and parent or guardian in the WBDR can be requested by parents or guardians by completing form F-40054A found at www.dhs.wisconsin.gov/cyshcn/birthdefects.

For additional information on birth defects in Wisconsin, refer to the website at www.dhs.wisconsin.gov/cyshcn/birthdefects. If you have questions, contact WBDR staff by email at dhswbdr@dhs.wisconsin.gov.

General information

Is this report a new report or a correction/addition to a previous report? ☐ New ☐ Correction

Have referrals been made to:

Children's Resource Center ☐ Yes ☐ No

Local Public Health Department ☐ Yes ☐ No

Birth to 3 Program ☐ Yes ☐ No

Other (specify): _____

Reporter information

Date of report: _____

Name and title of person completing form: _____

Phone of person completing form: _____

Email of person completing form: _____

Reporting source

Facility name: _____ Phone number: _____

Address: _____

Email: _____

Child's information

Child's name (Last, First, Middle): _____

Address – Street: _____

City: _____ State: _____ ZIP code: _____

Date of birth: _____ Medical record number: _____

Sex assigned at birth: ☐ Female ☐ Male ☐ Undesignated ☐ Ambiguous

Race

☐ American Indian or Native Alaskan☐ Asian☐ Black or African American☐ Native Hawaiian or other Pacific Islander☐ White☐ Other: _____☐ Unknown

Ethnicity

☐ Hispanic or Latino☐ Not Hispanic or Latino☐ Patient declined☐ Patient not asked☐ UnknownVital status: ☐ Alive ☐ Born alive, then died ☐ Stillborn >20 weeks gestation Date of death: _____

Birthweight

Grams: _____

Pounds and ounces: _____

Gestational age: _____ weeks (rounded to the nearest whole week)

Birth plurality: ☐ Single ☐ Twin ☐ Other multiple: _____If multiple, delivery order: ☐ First ☐ Second ☐ Other: _____Place of birth: ☐ Home birth ☐ Hospital ☐ Other: _____

Facility name: _____

Facility address: _____

Responsible party

Name (Last, First, Middle): _____

Address – Street: _____

City: _____ State: _____ ZIP code: _____

Preferred language: _____

Birth mother

Name (Last, First, Middle): _____

Address at time of pregnancy – Street: _____

City: _____ State: _____ ZIP code: _____

Date of birth: _____

Race

☐ American Indian or Native Alaskan☐ Asian☐ Black or African American☐ Native Hawaiian or other Pacific Islander☐ White☐ Other: _____☐ Unknown

Ethnicity

- ☐ Hispanic or Latino
☐ Not Hispanic or Latino
☐ Patient declined

- ☐ Patient not asked
☐ Unknown

Diagnostic and condition information

Diagnosing physician's name (Last, First): _____

Date of diagnosis: _____

Physician's specialty: _____

Diagnosing physician's facility: _____

Wisconsin Birth Defect Registry Reportable Conditions

Appendix A should be used to complete this section.

Code	Reportable condition	Prenatally diagnosed?		
		<input type="checkbox"/> Yes	<input type="checkbox"/> No	<input type="checkbox"/> Unknown
		<input type="checkbox"/> Yes	<input type="checkbox"/> No	<input type="checkbox"/> Unknown
		<input type="checkbox"/> Yes	<input type="checkbox"/> No	<input type="checkbox"/> Unknown
		<input type="checkbox"/> Yes	<input type="checkbox"/> No	<input type="checkbox"/> Unknown
		<input type="checkbox"/> Yes	<input type="checkbox"/> No	<input type="checkbox"/> Unknown

Child's primary care provider

Primary care provider's name (Last, First): _____

Name of clinic/hospital: _____

Provider's Specialty: _____ Phone number: _____

Maternal health history

Diabetes: ☐ Yes ☐ No ☐ Gestational only ☐ Unknown

Hypothyroidism: ☐ Yes ☐ No ☐ Gestational only ☐ Unknown

Other underlying illness(es): ☐ Yes ☐ No ☐ Unknown

If yes, specify: _____

Maternal phenylketonuria (PKU): ☐ Yes ☐ No ☐ Unknown

High blood pressure or hypertension: ☐ Yes ☐ No ☐ Unknown

Alcohol use: ☐ Yes ☐ No ☐ Unknown

Smoking: ☐ Yes ☐ No ☐ Unknown

Other substance: ☐ Yes ☐ No ☐ Unknown

If yes, specify: _____

Appendix A

Wisconsin Birth Defect Registry Reportable Conditions

ICD-10-CM Code	
A. Cardiovascular	
Q23.0	Aortic valve stenosis
Q21.1	Atrial septal defect
Q21.2	Atrioventricular septal defect
Q25.1	Coarctation of the aorta
Q20.1	Double outlet right ventricle
Q22.5	Ebstein's anomaly
Q23.4	Hypoplastic left heart
Q25.2, Q25.4	Interrupted aortic arch
Q25.21	Interruption of aortic arch
Q25.29	Other atresia of aorta
Q25.4	Other congenital malformations of aorta
Q25.40	Congenital malformations of aorta unspecified
Q25.41	Absence and aplasia of aorta
Q25.42	Congenital aneurysm of aorta
Q25.44	Congenital dilation of aorta
Q25.45	Double aortic arch
Q22.0, Q22.1	Pulmonary valve atresia and stenosis
Q22.0	Pulmonary valve atresia
Q22.1	Congenital pulmonary valve stenosis
Q20.4	Single ventricle
Q21.3	Tetralogy of Fallot
Q26.2	Total anomalous pulmonary venous return (TAPVR)
Q20.3, Q20.5	Transposition of the great vessels (arteries)
Q20.3	Transposition of great vessels
Q20.5	Corrected Transposition
Q22.4	Tricuspid valve atresia
Q20.0	Truncus arteriosus
Q21.0	Ventricular septal defect
B. Chromosomal	
Q90	Down syndrome
Q90.0	Trisomy 21, nonmosaicism
Q90.1	Trisomy 21, mosaicism
Q90.2	Trisomy 21, translocation
Q90.9	Down syndrome, unspecified
Q98.0, Q98.1, Q98.4	Klinefelter syndrome
Q98.0	Klinefelter syndrome karyotype 47, XXY
Q98.1	Klinefelter syndrome, male with more than two X chromosomes
Q98.4	Klinefelter syndrome, unspecified
Q91.4, Q91.5, Q91.6, Q91.7	Trisomy 13
Q91.4	Trisomy 13, nonmosaicism
Q91.5	Trisomy 13, mosaicism
Q91.6	Trisomy 13, translocation

Q91.7	Trisomy 13, unspecified
Q91.0, Q91.1, Q91.2, Q91.3	Trisomy 18
Q91.0	Trisomy 18, nonmosaicism
Q91.1	Trisomy 18, mosaicism
Q91.2	Trisomy 18, translocation
Q91.3	Trisomy 18, unspecified
Q96	Turner's syndrome
Q96.0	Karyotype 45, X
Q96.1	Karyotype 45, X iso (Xq)
Q96.2	Karyotype 46, X with abnormal sex chromosome, except iso (Xq)
Q96.3	Mosaicism, 45, X/46, XX or XY
Q96.4	Mosaicism, 45, X/other cell line(s) with abnormal sex chromosome
Q96.8	Other variants of Turner's syndrome
Q96.9	Turner's syndrome, unspecified
Q93.81	Velocardiofacial syndrome (Deletion 22q11.2)
C. Endocrine	
E03.0, E03.1	Hypothyroidism (congenital)
E03.0	Congenital hypothyroidism with diffuse goiter
E03.1	Congenital hypothyroidism without goiter
D. Eye	
Q12.0	Cataract (congenital)
Q12.2, Q13.0	Coloboma
Q12.2	Coloboma of lens
Q13.0	Coloboma of iris
Q11.0, Q11.1, Q11.2	Microphthalmia/anophthalmia
Q11.0	Cystic eyeball
Q11.1	Other anophthalmos
Q11.2	Microphthalmos
E. Gastrointestinal	
Q44.2, Q44.3	Biliary atresia
Q44.2	Atresia of bile ducts
Q44.3	Congenital stenosis and stricture of bile ducts
Q79.3	Gastroschisis
Q43.1	Hirschsprung disease
Q79.2	Omphalocele
Q40.0	Pyloric stenosis
Q42	Rectal/colonic atresia/stenosis
Q42.0	Congenital absence, atresia and stenosis of rectum with fistula
Q42.1	Congenital absence, atresia and stenosis of rectum without fistula
Q42.2	Congenital absence, atresia and stenosis of anus with fistula
Q42.3	Congenital absence, atresia and stenosis of anus without fistula
Q42.8	Congenital absence, atresia and stenosis of other parts of large intestine
Q42.9	Congenital absence, atresia and stenosis of other parts of large intestine part unspecified
Q41	Small bowel atresia/stenosis
Q41.0	Congenital absence, atresia and stenosis of duodenum

Q41.1	Congenital absence, atresia and stenosis of jejunum
Q41.2	Congenital absence, atresia and stenosis of ileum
Q41.8	Congenital absence, atresia and stenosis of ileum
Q41.9	Congenital absence, atresia and stenosis of small intestine, part unspecified
Q39.0, Q39.1, Q39.2, Q39.3,	Esophageal atresia/trachea-esophageal fistula
Q39.0	Atresia of esophagus without fistula
Q39.1	Atresia of esophagus with trachea-esophageal fistula
Q39.2	Congenital trachea-esophageal fistula without atresia
Q39.3	Congenital stenosis and stricture of esophagus
Q39.4	Esophageal web
F. Genitourinary	
Q56	Indeterminate sex and pseudohermaphroditism
Q56.0	Hermaphroditism
Q56.1	Male pseudohermaphroditism, not elsewhere classified
Q56.2	Female pseudohermaphroditism, not elsewhere classified
Q56.3	Pseudohermaphroditism, unspecified
Q56.4	Indeterminate sex, unspecified
Q64.0	Epispadias
Q64.10, Q64.19	Exstrophy of the bladder
Q64.10	Exstrophy of urinary bladder, unspecified
Q64.19	Other exstrophy of urinary bladder
Q64.12	Exstrophy of the cloaca
Q54 (excluding Q54.4)	Hypospadias
Q54.0	Hypospadias, balanic
Q54.1	Hypospadias, penile
Q54.2	Hypospadias, penoscrotal
Q54.3	Hypospadias, perineal
Q54.8	Other hypospadias
Q54.9	Hypospadias, unspecified
Q64.2	Posterior urethral valves (congenital)
Q60	Renal agenesis/hypoplasia
Q60.0	Renal agenesis, unilateral
Q60.1	Renal agenesis, bilateral
Q60.2	Renal agenesis, unspecified
Q60.3	Renal hypoplasia, unilateral
Q60.4	Renal hypoplasia, bilateral
Q60.5	Renal hypoplasia, unspecified
Q60.6	Potter's syndrome
Q64.3	Urethral atresia and stenosis
Q64.31	Congenital bladder neck obstruction
Q64.32	Congenital stricture of urethra
Q64.33	Congenital stricture of urinary meatus
Q64.39	Other atresia and stenosis of urethra and bladder neck
G. Hematologic	
D58.0	Hereditary spherocytosis
H. Musculoskeletal	

Q77.4	Achondroplasia
Q66.0, Q66.89	Clubfoot (congenital)
Q66.00	Congenital talipes equinovarus, unspecified foot
Q66.01	Congenital talipes equinovarus, right foot
Q66.02	Congenital talipes equinovarus, left foot
Q66.89	Other specified congenital deformities of feet
Q65	Hip dislocation (congenital)
Q65.0	Congenital dislocation of hip, unilateral
Q65.00	Congenital dislocation of unspecified hip, unilateral
Q65.01	Congenital dislocation of right hip, unilateral
Q65.02	Congenital dislocation of left hip, unilateral
Q65.1	Congenital dislocation of hip, bilateral
Q65.2	Congenital dislocation of hip, unspecified
Q65.3	Congenital partial dislocation of hip, unilateral
Q65.30	Congenital partial dislocation of unspecified hip, unilateral
Q65.31	Congenital partial dislocation of right hip, unilateral
Q65.32	Congenital partial dislocation of left hip, unilateral
Q65.4	Congenital partial dislocation of hip, bilateral
Q65.5	Congenital partial dislocation of hip, unspecified
Q65.6	Congenital unstable hip
Q65.8	Other congenital deformities of hip
Q65.81	Congenital coxa valga
Q65.82	Congenital coxa vara
Q65.89	Other specified congenital deformities of hip
Q65.9	Congenital deformity of hip, unspecified
Q78.0	Osteogenesis imperfecta
Q67.5, Q76.3, Q76.41	Scoliosis and kyphosis
Q67.5	Congenital scoliosis
Q76.3	Congenital scoliosis due to congenital bony malformation
Q76.41	Congenital kyphosis
Q76.411	Congenital Kyphosis, occipito-atlanto-axial region
Q76.412	Congenital kyphosis, cervical region
Q76.413	Congenital kyphosis, cervicothoracic region
Q76.414	Congenital kyphosis, thoracic region
Q76.415	Congenital kyphosis, thoracolumbar region
Q76.419	Congenital kyphosis, unspecified region
Q71, Q72, Q73	Limb deficiencies (reduction defects)
Q71.0-Q71.9	Reduction defects of upper limb
Q72.0-Q72.9	Reduction defects of lower limb
Q73.0-Q73.9	Reduction defects of unspecified limb
I. Neurologic	
Q00.0, Q00.1	Anencephaly
Q00.0	Anencephaly
Q00.1	Cranioarchischisis
Q01	Encephalocele
Q01.0	Frontal encephalocele
Q01.1	Nasofrontal encephalocele
Q01.2	Occipital encephalocele

Q01.8	Encephalocele of other sites
Q01.9	Encephalocele, unspecified
Q04.2	Holoprosencephaly
Q03	Hydrocephalus (congenital)
Q03.0	Malformations of aqueduct of Sylvius
Q03.1	Atresia of foramina of Magendie and Luschka
Q03.8	Other congenital hydrocephalus
Q03.9	Congenital hydrocephalus, unspecified
Q02	Microcephaly
Q04.6	Congenital cerebral cysts (porencephaly)
Q05, Q07.01, Q07.03	Spina bifida (without anencephalus)
Q05.0	Cervical spina bifida with hydrocephalus
Q05.1	Thoracic spina bifida with hydrocephalus
Q05.2	Lumbar spina bifida with hydrocephalus
Q05.3	Sacral spina bifida with hydrocephalus
Q05.4	Unspecified spina bifida with hydrocephalus
Q05.5	Cervical spina bifida without hydrocephalus
Q05.6	Thoracic spina bifida without hydrocephalus
Q05.7	Lumbar spina bifida without hydrocephalus
Q05.8	Sacral spina bifida without hydrocephalus
Q05.9	Spina bifida, unspecified
Q07.01	Arnold-Chiari syndrome with spina bifida
Q07.03	Arnold-Chiari syndrome with spina bifida and hydrocephalus
J. Orofacial	
Q30.0	Choanal atresia
Q37	Cleft lip with cleft palate
Q37.0	Cleft hard palate with bilateral cleft lip
Q37.1	Cleft hard palate with unilateral cleft lip
Q37.2	Cleft soft palate with bilateral cleft lip
Q37.3	Cleft soft palate with unilateral cleft lip
Q37.4	Cleft hard and soft palate with bilateral cleft lip
Q37.5	Cleft hard and soft palate with unilateral cleft lip
Q37.8	Unspecified cleft palate with bilateral cleft lip
Q37.9	Unspecified cleft palate with unilateral cleft lip
Q36	Cleft lip (without cleft palate)
Q36.0	Cleft lip, bilateral
Q36.1	Cleft lip, median
Q36.9	Cleft lip, unilateral
Q35	Cleft palate (without cleft lip)
Q35.1	Cleft hard palate
Q35.3	Cleft soft palate
Q35.5	Cleft hard palate with cleft soft palate
Q35.7	Cleft uvula
Q35.9	Cleft palate, unspecified
Q75.0	Craniosynostosis
Q16.0, Q17.2	Microtia/anotia
Q16.0	Congenital absence of (ear) auricle
Q17.2	Microtia

K. Pulmonary

E84	Cystic fibrosis
E84.0	Cystic fibrosis with pulmonary manifestations
E84.1	Cystic fibrosis with intestinal manifestations
E84.11	Meconium ileus in cystic fibrosis
E84.19	Cystic fibrosis with other intestinal manifestations
E84.8	Cystic fibrosis with other manifestations
E84.9	Cystic fibrosis, unspecified
Q79.0, Q79.1	Diaphragmatic hernia
Q79.0	Congenital diaphragmatic hernia
Q79.1	Other congenital malformations of diaphragm