Patient Label Here



Postpartum Child Encounter

Type of Birth: (Select one) □ Vaginal Birth □ Cesarean Birth

NEWBORN STATUS TAB	Birth Order: (select one)
NEWBORN STATUS TAB	
Was this baby admitted to this organization for Postpartum	•
Care only (the birth did not occur at the admitting hospital)?	
□ Yes □ No	Birth Weight:grams 🗆 Weight Unknow
If yes, complete all sections.	GA at Birth weeks days
If no, proceed to Section: BABY'S SEX	GA at Birthweeksdays
Admission date: dd / mm / yyyy Admission Time:	Baby's Sex: (select one) □ Male □ Female
Birth Location:	□ Ambiguous genitalia □ Unknown
□ Hospital and Name of Hospital:	
□ Home □ Nursing Station □ Other Ontario location	Arterial cord blood test status: (Select one) □ Done
□ Birth Centre & Name of Birth Centre:	□ Results pending □ Not done □ Unsatisfactory specimen
□ Outside of Ontario	□ Unknown
Newborn Transfer From:	Arterial Cord pH:
□ Hospital and Name of Hospital:	Autorial Coud Base France /defait.
□ Home Birth Midwifery (MW) Care and Name of MW Practice	Arterial Cord Base Excess/deficit:
Group:	Venous cord blood test status: (Select one) □ Done
□ Nursing Station	□ Results pending □ Not done □ Unsatisfactory specimen
□ Birth Centre and Name of Birth Centre:	□ Unknown
□ Other unit same hospital □ Other	2 OTKTOWT
Newborn Date of Birth: dd / mm / yyyy Time of Birth:	Venous Cord pH:
Torre of Births (C.) D. Varria al Birth D. Consume Birth	Venous Cord Base Excess/deficit:



What is newborn/infant's blood group and type,	SUMMARY TAB
ABO/Rh(D)? (Select one)	Bilirubin Measured Within 72 Hours Of Birth: (Select one) Yes - Transcutaneous bilirubin (TCB) Yes - Total Serum Bilirubin (TSB) No - Transferred Out/Discharged No - Declined No - Reason Unknown No - Reason Other Unknown
□ None □ Brachial plexus injury □ Cephalohematoma □ Clavicular Fracture □ Facial Nerve Injury □ Fracture – other □ Palsy-other	Hyperbilirubinemia Requiring Treatment: (Select one) □ Yes □ No □ Unknown
□ Other birth injury □ Unknown	Hyperbilirubinemia Treatment: (Select all that apply)
Neonatal Health Conditions: (select all that apply)	□ Phototherapy □ Treatment declined
□ None □ Failed CCHD screening □ Hypoglycemia	Highest Serum Bilirubin >340 umol/L:
□ NAS – Neonatal Abstinence Syndrome	□Yes □No □Unknown
□ Other □ Unknown	Highest Serum Bilirubin >425 umol/L
Newborn Congenital Anomalies Identified:	□ Yes □ No □ Unknown
□ None □ Suspected or Confirmed * Refer to addendum on the last page with pick list selections	Pain Relief Measures During First Blood Sampling by Heel Prick: (Select all that apply) □ Breastfeeding □ Skin to skin
Newborn Congenital Anomalies Suspected: (See Addendum)	□ Sucrose □ Other □ No pain relief measures □ No heel prick sampling
Newborn Congenital Anomalies Confirmed: (See Addendum)	□ Unknown if pain relief was provided



Neonatal/Infant Death: No Yes Yes, with termination of pregnancy *If yes, Neonatal/Infant Death Date: dd / mm / yyyyy Neonatal/Infant Death Time: hours / minutes	REASON FOR BREAST MILK SUBSTITUTE: Infant Medical: Hypoglycemia Inadequate Weight Gain Inborn Errors of Metabolism Significant weight loss in the presence of clinical indications Other clinical indications
Newborn Hearing Screen Result: (Select one)	Maternal Medical: □ Active herpes on breast □ Additional health concerns □ Contraindicated maternal medication □ HIV infection □ Severe maternal illness Other: □ Donor milk not available □ Informed Parent Decision to use Any Breast Milk Substitute □ Insufficient Maternal Milk Supply □ Birth mother not involved in care □ Not eligible for donor milk □ Unknown Consent for Use of Breast Milk Substitute: (Select one) □ Evidence that consent was obtained □ No evidence of consent □ Unknown





Neonatal Discharged or Iranster to: U Home
□ Transfer to NICU other hospital and Name of other hospital:
□ Transfer to NICU same hospital
□ Transfer to Paediatric unit same hospital
□ Transfer to other hospital and Name of other hospital:
□ Child and Family Services Apprehension
□ Transfer to other unit, same hospital
□ Other □ Unknown
Reason for Newborn Transfer:
Reason for Newborn Transfer:
Reason for Newborn Transfer: Requires higher level of care Other Unknown
Reason for Newborn Transfer: Requires higher level of care □ Other □ Unknown Neonatal Transfer Date: dd / mm / yyyy
Reason for Newborn Transfer: Requires higher level of care Other Unknown Neonatal Transfer Date: dd / mm / yyyy Neonatal Transfer Time:



ADDENDUM: Newborn Congenital Anomalies (Picklist Selections)

CENTRAL NERVOUS SYSTEM AND NEURAL TUBE DEFECTS

Absent cavum septum pellucidum (CSP) | Absent cerebellar vermis |Acrania or Anencephaly | Arachnoid cyst(s) | Arnold Chiari Malformation | Aqueductal stenosis | Corpus callosum – Agenesis (ACC) | Corpus callosum - Hypoplasia | Dandy walker malformation/variant (DWM) Encephalocele | Enlarged cisterna magna | Holoprosencephaly | Hydrocephalus | Hypotonia, unspecified | Lissencephaly | Macrocephaly | Microcephaly | Polymicrogyria | Posterior fossa cyst | Sacral agenesis | Sacral coccygeal teratoma (SCT) | Seizures | Spina bifida with hydrocephalus | Spina bifida without hydrocephalus Ventriculomegaly-Mild-Moderate (11-14.9 mm) | Ventriculomegaly-Severe (>15 mm) | Other – malformations of the nervous system | Other malformations of the brain

EYE ANOMALIES

Anophthalmos | Congenital cataract | Congenital glaucoma | Microphthalmos | Other- malformations of eye

EAR, FACE, AND NECK ANOMALIES

Ears - Anotia | Ears - Microtia | Choanal atresia | Macroglossia | Micrognathia | Nose - Absent | Nose - Hypoplastic | Retrognathia | Other - malformation of ear | Other - malformation of the face and neck

THORAX ANOMALIES

Bronchopulmonary sequestration (BPS) | Congenital high airway obstruction (CHAOS) | Cystic adenomatous malformation of lung (CCAM) | Diaphragmatic hernia (CDH) | Other – congenital malformations of lung | Other – malformations of the diaphragm

CARDIOVASCULAR ANOMALIES

Aortic arch – Double | Aortic arch – Interrupted | Aortic arch – Right | Aortic atresia/Hypoplastic aortic arch | Aortic valve stenosis | Arrhythmia | Atrial isomerism (heterotaxy) – left

| Atrial isomerism (heterotaxy) right | Atrial septal defect (ASD) | Atrioventricular septal defect (AVSD) Cardiomegaly | Coarctation of aorta | Complete/incomplete congenital heart block | Dextrocardia | Discordant atrioventricular connection | Double inlet ventricle (Single ventricle) Double outlet right ventricle (DORV) | Ebstein anomaly | Hypoplastic left heart (HLHS) | Hypoplastic right heart (HRHS) | Mitral valve atresia | Mitral valve insufficiency | Mitral valve stenosis | Patent ductus arteriosus (PDA) - >37 weeks | Patent/Persistent foramen ovale (PFO)/Premature closure of atrial septum | Pericardial effusion | Pulmonary valve atresia | Pulmonary valve dysplasia | Pulmonary valve stenosis | Situs inversus (cardiac and abdominal) | Tetralogy of Fallot (TOF) | Total anomalous pulmonary venous connection (TAPVC)/Partial anomalous pulmonary venous connection (PAPVC) | Transposition of great vessels (TGV) | Tricuspid atresia/ stenosis | Tricuspid regurgitation | Tricuspid valve dysplasia | Truncus arteriosus (common arterial truncus) | Vascular ring | Vena cava, bilateral



superior (SVC) | Vena cava, interrupted inferior (IVC) | Vena cava, persistent left superior (SVC) | Ventricular disproportion (RV/LV discrepancy) | Ventricular septal defect (VSD) | Other cardiac malformations

ORO-FACIAL CLEFTS

Cleft lip | Cleft palate | Cleft lip with cleft palate | Pierre Robin Sequence

GASTROINTESTINAL & ABDOMINAL ANOMALIES

Abnormal stomach (including small/ absent stomach) | Biliary atresia (atresia of bile ducts) | Bowel obstruction | Duodenal atresia/stenosis | Esophageal atresia (without fistula) | Esophageal with tracheoesophageal fistula (TEF) | Tracheoesophageal fistula (TEF) without esophageal atresia | Hirschsprung disease | Imperforate anus (congenital absence, atresia, stenosis of anus) | Large intestine atresia/stenosis | Pyloric stenosis | Rectal atresia/stenosis with/without fistula | Small bowel, abnormal | Small intestine atresia/stenosis (excluding duodenum) | Umbilical hernia | Other - malformations of gastrointestinal system

ABDOMINAL WALL DEFECTS

Gastroschisis | Omphalocele (exomphalos) | Other – congenital malformations of abdominal wall

URINARY ANOMALIES

Bladder/cloacal exstrophy | Congenital hydronephrosis | Cystic kidneys – other | Duplex kidney/collecting system | Echogenic kidney | Ectopic/pelvic kidney | Lower urinary tract obstruction | Megacystis | Megaureter | Multicystic dysplastic kidney(s) | Polycystic kidney, autosomal recessive | Polycystic kidney, autosomal dominant | Posterior urethral valves (PUV) | Prune belly | Renal agenesis, unilateral | Renal agenesis, bilateral | Renal cyst | Renal Dysplasia | Ureterocoele | Other – malformations of the urinary system

GENITAL ANOMALIES

Ambiguous genitalia/indeterminate sex | Cryptorchidism/undescended >37 weeks | Epispadias | Hydrocoele | Hypospadias | Other – malformations of female genitalia | Other – malformations of male genitalia

SKELETAL & LIMB ANOMALIES

Adactyly (absent fingers/toes) | Bowed/ curved long bone(s) | Club foot (talipes equinovarus) – bilateral | Club foot (talipes equinovarus) - unilateral Congenital hip dislocation/dysplasia | Craniosynostosis | Ectrodactyly (lobster-claw/cleft hand) | Hypotonia, unspecified | Limb reduction defect - upper limb | Limb reduction defect lower limb | Limb reduction defects of unspecified limb | Osteogenesis imperfecta | Polydactyly – hands Polydactyly – feet | Skeletal dysplasia | Syndactyly - hands | Syndactyly feet | Congenital malformations of the musculoskeletal system | Other - malformations of the spine & bony thorax (not including spina bifida) Other – malformations of the limb(s)

OTHER ANOMALIES/PATTERNS/ SYNDROMES

Congenital constriction bands/amniotic bands | Intrauterine growth restriction (IUGR) <10th %tile | Noonan syndrome | Oligohydramnios | Polyhydramnios | Pierre Robin Sequence | Potter's syndrome/sequence | Other – genetic syndrome



LYMPHATIC ANOMALIES & HYDROPS

Increased nuchal translucency (≥ 3.5 mm) | Cystic hygroma | Fetal ascites | Hydrops fetalis | Pleural effusion(s) (hydrothorax)

SKIN/HAIR/NAILS

Congenital ichthyosis | Cutis Aplasia | Epidermolysis Bullosa | Other – congenital malformations of skin | Other – congenital malformations of hair | Other – congenital malformations of nails

CHROMOSOMAL ANOMALIES

Down syndrome/Trisomy 21 | Patau syndrome/Trisomy 13 | Edwards syndrome/Trisomy 18 | Turner syndrome (45, X) | Klinefelter syndrome (47, XXY) | 47, XYY | Triple X syndrome (47, XXX) | Triploidy/polyploidy | 22q11.2 deletion syndrome/DiGeorge syndrome | Chromosome abnormality – other

TWINS

Twin-twin transfusion syndrome (TTTS) | Acardiac Twin (TRAP Sequence) |
Conjoined twins | Selective Intrauterine
Growth Restriction (sIUGR) | Twin
anemia polycythemia (TAPS) | Other
malformation(s) of twins

OTHER/UNKNOWN

Unknown | Other congenital malformations, not elsewhere classified