Twin embolisation syndrome—consequences in the living fetus: A short review

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Abstract
Twin embolization syndrome (TES) is a rare complication of monozygotic monochorionic twining following in utero demise of the co-twin and is related to the vascular connections in monochorionic placentas. Passage of thromboplastic materials, thrombus, debris and toxins from the dying twin’s circulation into the circulation of the surviving fetus results in ischemic structural defects of various systemic organs including the central nervous system (CNS). The purpose of this review is to analyse and present the consequences of TES in the living fetus.

Introduction
Twin embolization syndrome (TES) is a rare complication of monozygotic monochorionic twining following in utero demise of the co-twin and is related to the vascular connections in monochorionic placentas. Passage of thromboplastic materials, thrombus, debris and toxins from the dying twin’s circulation into the circulation of the surviving fetus results in ischemic structural defects of various systemic organs including the central nervous system (CNS) [1-3].

Monozygotic (MZ) or identical twins occur when a single ovum is fertilized to form one zygote which then divides into two separate embryos. By definition, monozygotic (MZ) twins carry an identical set of genetic information. About 1/3 of twin pregnancies is monozygotic pregnancies. The frequency of monozygotic twinning is almost the same among different countries and is not affected by mother’s age [4,5]. Monozygotic twinning starts by the stage of blastocyst at the end of the first week of pregnancy and is caused by the division of embryoblast in two embryonic structures. Eventually, two separate fetuses develop. So the two embryos, each one in a separate amniotic sac, grow into the same chorionic sac and share the same placenta, which is a monochorionic diamniotic placenta. In some cases, the primary separation of embryonic blastomeres causes monozygotic diamniotic twin pregnancies with two separate placentas [5].

Monozygotic twins occur in 3-5 per 1000 pregnancies. Monozygotic twins can be monochorionic or dichorionic. Approximately 75% of monozygotic twins are monochorionic. Only monochorionic twins are at risk for TES in the surviving fetus. TTS occurs in 5-38% of monochorionic twins. Thus, TTS only occurs in same sex, monozygotic twins with monochorionic placentation. Twin embolization syndrome is the most severe complication of TTS, so it is rarer than TTS.

Twin embolization syndrome is thought to result from the passage of thromboplastic material, thrombus, debris and toxins into the circulation of the surviving twin, which leads to emboli, thrombosis, clots and circulatory problems, that cause ischemic structural defects in various organs (particularly the highly vascularised organs such as the central nervous system, gastrointestinal tract, and genitourinary system). The nature of damage to the surviving fetus appears to be related to the gestational age at the time of death of the co-twin (demise in early pregnancy results in atresia and tissue loss; demise later in pregnancy results in tissue infarction). These thromboplastic agents precipitate disseminated intravascular coagulation in the surviving fetus, with a hypercoagulable state due to a relative fetal antithrombin III deficiency. Acute haemodynamic shift from live to dead fetus resulting in hypoperfusion is more recently thought to play a role. In conclusion, anomalies of central nervous system and other systems appear in the surviving fetus and end up to twin embolization syndrome.

Materials and methods
Many medical papers have presented implications of TES in the surviving fetus as analysed below.

Results
CNS anomalies
Fetal ventriculomegaly
Fetal ventriculomegaly refers to the presence of dilated cerebral ventricles in utero. Fetal ventriculomegaly (ventricle width >10 mm) is an important finding in itself and it is also associated with other central nervous system abnormalities. The estimated prevalence may be at about 0.9% of all pregnancies, with a slightly increased male predilection [6].

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Fetal ventriculomegaly is defined as:

- >10 mm across the atria of the posterior or anterior horn of lateral ventricles at any point in the gestation
- alternatively, a separation of more than 3 mm of the choroid plexus from the medial wall of the lateral ventricle may be used

The severity of ventriculomegaly can be further classified as:

- mild/borderline fetal ventriculomegaly: lateral ventricular diameter between 10-12 mm
- moderate fetal ventriculomegaly: 12.1-15 mm
- fetal ventriculomegaly (also sometimes classified as fetal hydrocephalus): severe lateral ventricular diameter >15 mm.

Pathology

Development of lateral ventricles differs depending on the trimester in which ventriculomegaly was created.

First trimester

The choroid plexus regularly fills the entire lateral ventricle, bilaterally.

Second trimester

- The choroid plexus begins to recede posteriorly but remains in close contact with the medial and lateral walls of the bodies and atria of the ventricles.
- Likewise, the lateral cerebral ventricle is large relative to the cerebral hemispheric width [7].

Clinical presentation

Infants with mild self-limited ventriculomegaly usually don’t have any symptoms. If the ventriculomegaly is progressive the baby may show the signs and symptoms of hydrocephalus once it is born.

An infant with hydrocephalus may have:

- abnormally rapid head growth
- abnormally full fontanel
- distended scalp veins
- eyes that cannot look upward or appear to be staring downward (sunset sign)
- developmental delays
- irritability or abnormal sleepiness
- poor feeding
- vomiting

Diagnosis

The findings must be carefully evaluated, because of the association of fetal ventriculomegaly with other central nervous system anomalies. Pseudo-hydrocephalus as a possible diagnosis must be excluded and ultrasound is the screening modality of choice for initial evaluation. Fetal brain MRI may be useful too [7].

Porencephaly

Porencephaly is a rare congenital disorder that results in cystic degeneration and encephalomalacia and the formation of porencephalic cysts. The term is used variably among radiologists with its broadest definition being a cleft or cystic cavity within the brain, and its more narrow definition being a focal cystic area of encephalomalacia that communicates with the ventricular system and/or the subarachnoid space. Generally this term is used to describe any fluid-filled cavity in the fetal or neonatal brain [7].

Pathology

Porencephalic cysts are uncommon, but in some cases of TES severe porencephaly subsequently developed. The cysts are typically lined by white matter. They are thought to occur from focal encephalomalacia due to a localized cerebral insult most frequently during early gestation. Gliosis will develop if the insult is late enough, usually thought to be after the start of the third trimester, although perhaps insults as early as 20 weeks of gestation may result in gliosis [7].

Clinical presentation

Clinical manifestations are extremely variable. Patients may be asymptomatic or in other cases they develop spasticity, seizures, language impairment, mental retardation and motor deficits in the first year of life [6].

Diagnosis

Porencephalic cysts may appear as one or more intracranial cysts, on antenatal ultrasound scan, that communicate with the ventricular system and subarachnoid space. There may also be displacement of the midline ventricular echo because of asymmetrical ventricles [7].

The cysts’ border is well-defined, on a CT scan, and occasionally enlarging cysts result in local mass effect [7]. On a MRI scan, the cyst is lined by white matter which may or may not demonstrate evidence of gliosis. Importantly the cyst is not lined by grey matter, helpful in distinguishing them from arachnoid cysts and schizencephaly [7]. Head circumference is variable. It may be normal or small, or alternatively synechiae can create a one-way valve effect with progressive enlargement of the cyst and skull expansion or hydrocephalus, resulting in an enlarged head.

Fetal cerebral atrophy

Fetal brain atrophy or shrinkage is rare and the cause is unknown in most cases. Among the most common risk factors is maternal alcohol consumption and intrauterine infection (Cytomegalovirus).

Pathology

Passage of thromboplastic materials, thrombus, debris and toxins from the dying twin's circulation into the circulation of the surviving fetus results in ischemic structural defects of various systemic organs including the central nervous system (CNS). Multiple lines of evidence support that cerebral ischemia is often the major factor that initiates cerebral injury in VLBW infants. In the near-term fetus, even brief severe global asphyxia, caused by a 10-minute occlusion of the umbilical cord, can result in prolonged hypoperfusion to cerebral gray and white matter. Systemic hypotension arising from intermittent or partial umbilical cord occlusion produced a variable degree of WMI in addition to primary damage to the cerebral cortex. Hence, cerebral hypoperfusion in conjunction with hypoxia appears to be a critical factor to generate significant preterm WMI. The spectrum of white matter pathology includes three major identifiable forms: focal cystic necrosis, focal microscopic necrosis and diffuse non-necrotic lesions.
Clinical presentation

The neonate can be small for gestational age, and can have bulging anterior and posterior fontanels, flat mid-face, smooth philtrum, hypertonia and increased deep tendon reflex in all extremities. The infant may have normal feeding tolerance, but can develop frequent neonatal seizures. 1 month after birth, the infant appears with suboptimal neurodevelopment and neurological signs of hypertonia, spastic tone, no eye contact and no social smile [9].

Diagnosis

On an ultrasound scan we may see decreased amniotic fluid and abnormal structures of the brain. Most of the space of the anterior fossa can be occupied by anechoic fluid or marked dilatation of the subarachnoid space around the brain, leading to a wide distance from the cranium to the brain surface. The cerebral hemispheres might display shrinkage to the center of the skull as a thin covering of the mid-brain. The thalami, brainstem, cerebelli and other structures in the posterior fossa can be well preserved and relatively normal in size.

On color-flow imaging, the circle of Willis can be visualized clearly and spectral Doppler ultrasonography may demonstrate a relatively low peak systolic velocity in the middle cerebral artery [9]. It is considered magnetic resonance imaging (MRI) of the fetal brain to be effective in confirming or denying diagnosis of fetal cerebral defects when ultrasonography was inconclusive or incomplete. Also, histological studies can confirm or deny ultrasonographic evidence [10].

Cystic encephalomalacia

The terms “Cystic encephalomalacia” is used to describe the presence of areas of necrosis that develop into cystic lesions inside the brain. These lesions are generally due to severe asphyxia and/or hypotension [11,12].

Pathology

Encephalomalacia, from the Greek “brain softening,” refers to diffuse cerebral parenchymal volume loss. This typically occurs between the 20th and 30th week of gestation with the most common etiologies being maternal infection, trauma, or vascular insult. Impaired brain perfusion leads to neuronal injury and cell death. Neuronal cell loss occurs in two phases: primary and delayed. The high rates of cellular metabolism in the developing fetal brain make neurons particularly sensitive to ischemia and lead to primary neuronal cell loss within 30 minutes of onset of hypoxia and ischemia. However, many neurons are spared during the initial insult. Delayed loss results from free radical production. The cytotoxicity-mediated apoptosis cascade usually occurs in the days following the insult. In contrast to porencephaly, or focal cystic white matter defects communicating with the ventricular system, encephalomalacia results in a global destruction of brain matter with symmetric bilateral effects. Because of neuronal cell death, diffuse brain atrophy results. In uterus and postpartum outcomes are variable. However, this type of injury generally confers poor prognosis with neurodevelopmental delay, seizure disorders, and sensory and motor deficits [13].

Clinical presentation

The infant may have hypotonia during the first days of life. At 1 month of life the infant might still have hypotonia and delayed development and also it might be lethargic and irritable. At 3 months of life hypertonia and hyperreflexia are possible for the infant and it may appear still as irritable [8,13].
deviations (SDs) below the average. The measurement value also may be designated as less than the 3rd percentile [15].

Gastrointestinal anomalies

Gastrochisis

Gastrochisis refers to extra-abdominal herniation (eversion) of fetal or neonatal bowel loops (and occasionally portions or the stomach and or liver) into the amniotic cavity through a para-umbilical abdominal wall defect. The estimated incidence is at around 1-6 per 10,000 live births. There may be a male predilection and an increased incidence with younger maternal age.

Pathology

This condition occurs when an opening forms in the baby's abdominal wall. The baby's bowel pushes through this hole. The bowel then develops outside of the baby's body in the amniotic fluid. The passage of thromboplastic materials as it happens in TES Syndrome can be a cause, because this syndrome causes problems in vascular supply to the area in the abdominal wall adjacent to the umbilicus. This anomaly does not have a surrounding membrane (unlike an uncomplicated omphalocoele). It is the small bowel that herniates most often. The defect is invariably on the right side and usually measures between 2-4 cm. There is no covering membrane or membrane remnant [16].

Clinical presentation

Half of malformations are considered related to the gastrochisis (intestinal atresia or stenosis, malrotation, cryptorchidism, amyoplasia, urinary tract obstruction). Other associated malformations occur which are not recognized to be secondary to the gastrochisis. Maternal serum alphafetoprotein is increased. Prominent among these are cardiac and limb defects. Fetal and neonatal mortality are increased, but neither appear related to lethal malformations [17].

Diagnosis

It is possible for gastrochisis to be detected in the third month of pregnancy. However, doctors most often perform evaluations for it at 20-24 weeks, after it has shown up on an ultrasound. It is most commonly diagnosed by ultrasound around weeks 18-20 of pregnancy.

An evaluation for gastrochisis consists:

- An ultrasound
- Possibly an MRI and/or a fetal echocardiogram to test baby's heart function
- Maternal serum alphafetoprotein measurement
- A meeting with a nurse, social worker and genetic counselor
- A team meeting with a maternal-fetal medicine specialist (MFM), pediatric surgeon and neonatologist

An important part of the evaluation is determining whether the condition is gastrochisis or omphalocoele. These conditions can sometimes look similar on an ultrasound. In omphalocoele, a sac from the umbilical cord covers and protects the intestines that are outside of the baby's body [16].

Small bowel atresia

Small bowel atresia can be ileal atresia, which is a congenital abnormality where there is significant stenosis or complete absence of a portion of the ileum, or jejunal atresia, which is a congenital anomaly characterised by obliteration of the lumen of the jejunum. The site of the atresia can be anywhere from the ligament of Treitz to the jejunoileal junction. There can be more than one atretic segment.

Pathology

Ileal atresia results from a vascular accident in utero that leads to decreased intestinal perfusion and subsequent ischemia a segment of bowel, or neonatal intestinal obstruction. As far as TES Syndrome is concerned, thromboplastic materials that pass to the circulation of the living fetus, cause clots and also decreased intestinal perfusion. This leads to narrowing, or in the most severe cases, complete obliteration of the intestinal lumen.

Clinical presentation

Common clinical characteristics of patients with jejunoileal atresia include the following:

- Polyhydramnios on antenatal ultrasonography (28%)
- Prematurity (35%)
- Low birth weight (25-50%)

Classic signs include the following:

- Bilious emesis that warrants emergency surgical evaluation (most patients)
- Abdominal distention (in distal atresia)
- Jaundice (32%)
- Failure to pass meconium in the first 24 hours (rule out Hirschsprung disease; passage of meconium does not rule out intestinal atresia)

Signs of continuous fluid loss include the following:

- Dehydration, manifested by sunken fontanel and dry membranes
- Decreased urine output (the best clinical indication of tissue perfusion)
- Tachycardia
- Decreased pulse pressure
- Low-grade fever
- Neurologic involvement, manifested by irritability, lethargy, or coma [18]

Diagnosis

Jejunoileal atresia can be identified on the basis of polyhydramnios present during prenatal ultrasonographic evaluation, bilious vomiting, abdominal distention, and jaundice. Some patients may not pass meconium in the first day of life.

Plain radiographs of the abdomen typically show the classic double-bubble sign: two distinct gas collections or air-fluid levels in the upper abdomen, resulting from the markedly dilated stomach and proximal duodenal bulb. There can be multiple dilated small bowel loops proximal to the atresia and the number of dilated loops increase as point of atresia becomes more distal.

In fluoroscopy, Contrast enema typically shows micro colon (small unused colon).
Antenatal ultrasound may show:
- dilated proximal bowel loops, often greater than 7 mm
- evidence of an in utero bowel perforation
- polyhydramnios, especially in cases where the atresia is proximal

In any case, small intestinal atresia must be differentiated from malrotation with midgut volvulus, total colonic Hirschsprung disease and meconium ileus [18].

**Urogenital Anomalies**

**Fetal renal cortical necrosis**

Renal cortical necrosis is a rare cause of acute renal failure secondary to ischemic necrosis of the renal cortex. Renal cortical necrosis is usually extensive, although focal and localized forms occur. In most cases, the medulla, juxtamedullary cortex, and a thin rim of subcapsular cortex are spared.

Renal cortical necrosis is classified into 5 pathologic forms, depending on severity, as follows:

1. Focal pathologic form - Kidneys show focally necrotic glomeruli without thrombosis and patchy necrosis of tubules
2. Minor pathologic form - Larger foci of necrosis are evident with vascular and glomerular thrombi
3. Patchy pathologic form - Patches of necrosis may occupy two thirds of the cortex
4. Gross pathologic form - Almost the entire cortex is involved; thrombosis of the arteries is more widespread
5. Confluent pathologic form - Kidneys show widespread glomerular and tubular necrosis with no arterial involvement

Renal cortical necrosis accounts more than 20% of acute renal failure during the third trimester of pregnancy. Renal cortical necrosis was detected by postmortem examination in 5% of infants aged 3 months or younger at death [19].

**Pathology**

Although the pathogenesis of the disease remains unclear, the presumed initiating factor is intense vasospasm of the small vessels. If this vasospasm is brief and vascular flow is reestablished, acute tubular necrosis results. More prolonged vasospasm can cause necrosis and thrombosis of the distal arterioles and glomeruli, and renal cortical necrosis ensues. In hemolytic-uremic syndrome (HUS) and septic abortion, an additional mechanism involves endotoxin-mediated endothelial damage that leads to vascular thrombosis. Studies have shown that patients with HUS with thrombotic microangiopathy (TMA) involving arteries have a higher likelihood of progressing into acute cortical necrosis compared with patients with predominant glomerular TMA. Renal cortical necrosis in placental abruption may be due to a combination of a hypercoagulable state, endothelial injury, and intravascular thrombosis.

In TES Syndrome, because of the passage of thromboplastic materials and toxins into the circulation of the surviving fetus, the perfusion to kidneys is low and this leads to ischemia and fetal renal cortical necrosis. Other causes of this condition are congenital heart disease, fetal-maternal transfusion, dehydration, perinatal asphyxia, anemia, placental hemorrhage, severe hemolytic disease and sepsis [18].

**Clinical presentation**

Kidney findings may include abdominal or bilateral costovertebral tenderness and palpable, tender kidneys.

**Symptoms of Renal Cortical Necrosis (RCN)**
- Flank Pain
- Hematuria
- Proteinuria
- Lower Urine Output
- Water Retention-
- Edema feet (swelling of feet)
- Enlarged liver
- Pleural effusion (fluid collection around lungs) [19,20]

**Diagnosis**

- Serum Electrolytes- High potassium level
- CBC- Low hemoglobin and red blood cell count
- Coagulation Panel- Blood clot disorder
- Urinalysis- Blood and protein in urine
- Ultrasound- Kidney size is smaller
- CT scan of the kidneys
- Kidney Biopsy- shows necrosis and blood clots [20]

**Respiratory Anomalies**

**Fetal hydrothorax**

A fetal hydrothorax refers to fluid in the fetal thoracic cavity. In many cases it represents a fetal pleural effusion. In selected cases it can be treated by in utero thoracocentesis or a formation of a in utero pleuro-amniotic shunt [7].

**Pathology**

The etiology of congenital hydrothorax is highly variable. Antenatal primary hydrothorax typically corresponds to congenital chylothorax, and is mainly due to lymphatic leakage because of a malformation, atresia or fistula, in the development of pulmonary lymphatic vessels of the thoracic duct. As far as TES Syndrome is concerned, the pulmonary lymphatic vessels have an obstruction or a malfunction due to passage of thromboplastic materials of dead fetus into the circulation of the surviving fetus. This leads to lymphatic leakage and chylothorax [21].

**Clinical presentation**

Primary fetal hydrothorax can resolve spontaneously in utero. However, there are specific signs that suggest the presence of an elevated intrathoracic pressure and therefore an increased risk for developing hydrops. Some of these signs are enlargement of the pleural effusion, evident mediastinal shift, and increased cardiac preload (abnormal ductus venosus blood flow, tricuspid regurgitation, and cardiomegaly). Also, hydramnios secondary to enlarged effusions causing esophageal compression and impeding fetal swallowing might lead to a poorer prognosis due to premature labor. The presence of any these markers, or the actual onset of hydrops, would be indications for urgent fetal intervention.
Diagnosis

Prenatal diagnosis of congenital hydrothorax is based on the demonstration of fluid collection in the thorax, surrounding the lungs uni or bilaterally, except for the hilum. It is usually diagnosed on the four-chamber view. If unilateral, it is considered as hypertensive if the intrathoracic pressure is high and the heart and all viscera are pushed into the contralateral hemithorax. If bilateral, it may produce increased venous return pressure and secondary hydrops [1,2,7,11]. Regarding the etiologic diagnosis, an initial thoracocentesis is useful for diagnosing congenital chylothorax (more than 80% of lymphocytes in the blood count) and for decompressing the lungs [12,13]. Usually, this procedure alone will not resolve the problem, since in more than 70-80% of the cases the liquid will reaccumulate quickly.

Differential diagnosis with pericardial effusion is usually easy and straightforward, whenever the scan is performed by a medium experienced examiner. In pericardial effusion the fluid is surrounding the heart and both lungs remain attached to the posterior wall of the chest [21].

Diagnosis

Prenatal: Sonographic evaluation of the fetus is commonly performed during pregnancy in order to diagnose any abnormalities that may occur through the first, second or third semester of it. As far as monochorionic twin pregnancies are concerned, sonography is a method capable of diagnosing demise of one of the twins in utero (vanishing twin syndrome), with or without associated bleeding referred from the mother [8]. The presence of a dead twin combined with a surviving twin with various anomalies may suggest the diagnosis of the twin embolisation syndrome. Sonographic evaluation of the surviving twin may diagnose CNS anomalies such as fetal ventriculomegaly, porencephaly, fetal cerebral atrophy, cystic encephalomalacia and microcephaly. Non CNS anomalies such as small bowel atresia, fetal hydrothorax, gastrochisis, aplasia cutis, hydrothorax and renal cortical necrosis may also be diagnosed through sonography. Twin embolisation syndrome is not usually diagnosed in cases that the death of the co-twin happened during the first semester of the pregnancy [10,11].

Another method that is used to diagnose twin embolisation syndrome is Doppler ultrasound. Doppler assessment of the placental circulation plays an important role in screening for impaired placenta and its complications of pre-eclampsia, intrauterine growth restriction and perinatal death. Assessment of fetal circulation is essential in understanding the pathophysiology of a wide range of pathological pregnancies and their clinical management. In Doppler ultrasound scanners, a series of pulses is transmitted to detect movement of blood. It is proved that color Doppler ultrasound sonography can play an important role in identifying the communicating placental vessels in pregnancies with twin-to-twin transfusion syndrome or its severe form, the twin embolization syndrome [12].

Postnatal: After birth, the surviving twin who has CNS and non-CNS anomalies, which are referred in previous paragraphs, has to be put through some more diagnostic examinations. Magnetic resonance imaging (MRI) is a powerful and versatile diagnostic tool. Applications of MR technology are rapidly expanding for all patient populations, including infants receiving newborn intensive care. The surviving infant now routinely undergoes MRI of the head, chest, abdomen, spine and pelvis for a multitude of diagnostic purposes. MRI techniques are superior to most other diagnostic and prognostic neuroimaging applications. X-rays or computed tomography (CT) scans can also be used but, unlike MRI, they use ionizing radiation. Instead, MRI uses an extremely powerful static magnetic field, rapidly changing gradient magnetic fields, and radiofrequency electromagnetic impulses to produce detailed anatomic or functional images of the brain and other soft tissues of the body of the surviving twin [14].

After the demise of one of the twins during pregnancy, maternal and fetal monitoring is performed until birth in order to prevent complications in the surviving twin.

Maternal monitoring includes:
- Routine antenatal investigations (ABO Rh, Hb, GCT, Urine routine and microscopy and viral markers)
- Weekly coagulation profile (platelet count, PT, aPTT, BT and CT)
- FDP and D-dimer
- Fetal monitoring includes:
  - Daily fetal movement count
  - Biweekly NST in pregnancies more than 32 weeks
  - Biweekly USG with biophysical profile and colour Doppler [13]

Theoretically, premature delivery and treatment of the surviving fetus at an early stage may reduce the degree of cerebral as well as other organ damage. The benefits of this theoretical approach must be balanced by the inevitable risks of prematurity, complications of anticoagulant therapy and risks after treatment such as fetoscopic laser vessel occlusion [2].

Discussion

In cases of severe twin-to-twin transfusion syndrome, may not only one but both of the co-twins die prenatally. The most severe form of twin-to-twin transfusion syndrome with acute hydramnios and stuck twin usually occurs between 18 and 26 weeks gestation, with a survival rate of 20-45% for those identified before 28 weeks. Passage of thromboplastic-like material or embolic debris to the circulation of the surviving twin may cause multiple organ ischemia which has poor prognosis [2]. Apart from this, the surviving twin is possible to face premature birth (35.1 -38.2 week) or very premature birth (< 32 week) which increases the risk of mortality. Low birth weight is also linked to twin embolisation syndrome and affects the prognosis [15].

The surviving twin will also have associated with the twin embolisation syndrome congenital anomalies which will affect prognosis. Three common anomalies are ventriculomegaly, porencephaly and microcephaly. Cases of ventriculomegaly resolved, regressed or remained stable in utero will exhibited good prognosis (85% of all cases). Ventriculomegaly with an transverse atrial size >= 12 mm or progression in utero are usually associated with a poor prognosis, which should be observed carefully [16]. As for porencephaly, the extent of impairment is somewhat related to the size and location of the lesion [17]. Microcephaly generally has good prognosis but can lead to disability [18].

When the diagnosis of an intrauterine fetal death, in the setting of a monochorionic gestation, is made near term, immediate delivery seems appropriate to improve the prognosis of the surviving twin. Earlier in pregnancy, in severe previable twin embolisation syndrome, fetoscopic laser occlusion of placental vessels or cord occlusion of the dead twin was reported to prevent further damage to the surviving twin. Meticulous sonographic evaluation permits counseling for the appropriate treatment, prognosis and risks in twin embolisation syndrome [2].
Case studies

A 19 year old primigravida was referred for sonographic evaluation at 14 weeks of her first, spontaneous pregnancy. Vaginal sonography revealed a twin pregnancy. One fetus had no heartbeat, while the other fetus had a normal sonographic examination. Both fetuses had CRL and BPD measurements appropriate for gestational age. A thin dividing membrane was clearly seen, revealing a monochorionic, diamniotic twin pregnancy with missed twin.

Owing to religious reservations, the triple test wasn’t performed. At 22 weeks gestation the patient was referred for another sonographic examination. The surviving fetus was found to have bilateral ventriculomegaly (13 mm). The third and fourth ventricles were normal. The nuchal fold, heart, skin and thorax were found to be normal. Bowel distension suggesting obstruction was observed with multiple intraabdominal calcifications. AC, HC and FC had dimensions compatible with a gestation of two weeks earlier, with a normal amount of amniotic fluid suggesting early intrauterine growth retardation. The second fetus, of a size compatible with a gestation of 12 to 13 weeks, was clearly in an individual sac with no pulse, located above the cervix. Cordogenesis or amnioncensis was denied by the couple for religious reasons.

At 30 weeks gestation sonographic evaluation of the fetal brain revealed right-sided ventriculomegaly of 11.4 mm and left-sided unilateral porencephaly. The size of the penis was small. BDL, FL and AC were compatible to 28 weeks and the estimated fetal weight was 1210 g. The second, missed twin, had a CRL of 45 mm. The result of Doppler flow studies performed on the umbilical artery and the carotid artery of the surviving twin were normal.

At 35 weeks gestation the patient experienced premature contractions and minimal vaginal bleeding. Clinical examination revealed that the uterus was distended, suggesting polyhydramnios, with no cervical dilatation. Sonographic examination verified severe polyhydramnios. Asymmetrical hydrocephalus, with porencephaly of the right cortex and left posterior cortex, was present, and the fetal abdominal sonography revealed bowel dilatation with multiple calcifications. A cyst measuring 15mm in diameter was found in the spleen, probably secondary to an infect.

At 36 weeks gestation the patient had a normal delivery of a boy with Apgar scores of 5 and 9 at 1 and 5min. After tracheal suction the newborn was stable with Apgar scores of 5 and 9 at 1 and 5 min. Blood pressure of 60/37 mm Hg.

The placenta was found to have bilateral ventriculomegaly due to progressive porencephaly with minimal brain tissue. The karyotype was normal (46 X,Y). Parenteral nutrition was started and a nasogastric tube was used to drain excessive secretions. However, the infant died after 2 months owing to aspiration pneumonia. Autopsy was denied by the parents for religious reasons [2].

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