The EXIT (ex-utero intrapartum treatment) procedure – from the paediatric ENT perspective

Procedura EXIT (ex-utero intrapartum treatment) – prospettive otorinolaringoiatriche pediatriche

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SUMMARY
The main principle of the EXIT procedure is to maintain uteroplacental circulation with neonatal anaesthesia by controlled uterine hypotonia. This enables securing the foetal airways and decompress or resect large neck and mediastinal foetal masses. The authors present their experience with use of the EXIT procedure in 7 foetuses in whom evaluation and management of the airways were performed. In 4 patients, the neck mass was surgically removed in the neonatal period, in 1 the propranolol treatment was introduced. Two newborns died shortly after the EXIT procedure. The EXIT procedure allows the paediatric otolaryngologist to provide airway patency of newborns during delivery. Both ultrasound and MR imaging are crucial in the prenatal assessment of foetal head and neck masses. Their application in the evaluation of any foetal anomaly is essential for proper prognosis and treatment. Maternal monitoring for complications such as polyhydramnios and preterm labour are important in planning and desirability of the EXIT procedure.

KEY WORDS: EXIT procedure • Foetal neck masses • Foetal airways • Prenatal imaging • Teratoma • Lymphatic malformation

INTRODUCTION
The EXIT (ex-utero intrapartum treatment) procedure was initially used for reversing tracheal occlusion in foetuses with CDH (Congenital Diaphragmatic Hernia). The main principle of the EXIT is to maintain uteroplacental circulation with neonatal anaesthesia by controlled uterine hypotonia. This enables to secure the foetal airways and decompress or resect large neck and mediastinal foetal masses and provide vascular access or cannulation for the ECMO. Allowing the paediatric otolaryngologist to gain adequate control of the airway can thus improve the chances of survival in such cases. According to Hedrick, the indications for EXIT can be based on the goal of the procedure and at the same time include the type of lesion and diagnosis. The EXIT to airway procedure requires the presence of a paediatric otolaryngologist and is attempts to evaluate and secure the airways during delivery because of the presence of a congenital neck mass with airway obstruction or CHAOS syndrome, or the necessity for reversal of tracheal occlusion. The EXIT to resection procedure involves the partial or complete surgical removal of the congenital head and neck masses. The other indications are: the EXIT to ECMO in congenital heart anomalies and the EXIT to separation procedure (in conjoined twins). In contrary to the caesarean section, during the EXIT procedure the achievement of tocolysis is the main aim to provide placental support. After hysterotomy, the foetal
head and neck and one arm are delivered and the rest of the foetal body and umbilical cord remain in the uterus to preserve heat and fluid loss.

The authors use the airway algorithm to secure the airways as presented by Marwan et al. from Cincinnati Children’s Hospital. First, direct laryngoscopy is performed. If not achieved, the next step is to intubate the foetus under control of a rigid bronchoscope with a diameter of 2.5-3.0 mm. In some cases, partial surgical reduction in the tumour mass is necessary. If the intubation is not possible, the ENT specialist performs tracheostomy. After securing the airway, surfactant is administered prophylactically to prevent barotrauma of the lungs and respiratory failure. The newborn is ventilated by the Ambu bag, and before cutting the umbilical cord, the ENT team checks the position of the tracheostomy tube using a flexible bronchoscope.

Clinical cases

The authors present their experience with use of the EXIT procedure in management of 7 foetuses in whom the evaluation and management of the airways were performed. In 4 patients, the neck mass was surgically removed in the neonatal period and in one propranolol treatment was introduced. Two newborns died shortly after the EXIT procedure.

Case 1. Reversal of foetal tracheal occlusion

A female foetus with severe CDH and pulmonary hypoplasia underwent reversal tracheal occlusion with the PLUG (Plug the Lung Until it Grows) balloon (the FETO procedure was performed in London – “foetal endoscopic surgery”) at a gestational age of 26 weeks. Emergency delivery by the EXIT procedure was performed at 32 weeks of gestation because of unrestrained systolic uterus function and heart rate deceleration. During rigid bronchoscopy, the balloon was removed and the neonate was intubated. The chest X-ray revealed left lung agenesis. The baby died on the 2nd day after birth for severe respiratory deficiency. Postmortem examination confirmed left pulmonary agenesis and right lung hypoplasia due to lack of efficacy of the PLUG balloon.

Case 2. Congenital High Airway Obstruction Syndrome (CHAOS)

A female foetus was prenatally diagnosed with laryngeal atresia and polyhydramnios. Parents sought the centre in which the child could be born and undergo laryngeal reconstruction surgery. They obtained the consent of the Polish Ministry of Health to fund the labour, the EXIT procedure and laryngeal reconstruction surgery in the US. The only requirement was to maintain pregnancy to 32 weeks. Unfortunately, spontaneous delivery started at gestational week 27 with premature rupture of membranes. The emergency EXIT was performed with tracheostomy attempt. The foetus presented with hydrops fetalis and died shortly after the EXIT procedure. Postmortem examination revealed complete laryngeal and tracheal agenesis.
Cases 3-7. Foetal neck masses

The most common indication for the EXIT procedure in our patients was foetal neck mass, most frequently lymphatic malformations and teratomas. In such cases, prenatal imaging of the neck mass related to airway structures and oesophagus is essential to optimise foetal outcome. In 5 foetuses (4 boys and 1 girl), the presence of a neck mass was prenatally diagnosed with ultrasound and MR imaging. It revealed in 2 cases lymphatic malformation, in another 2 teratoma and in 1 infantile haemangioma. In all cases the EXIT procedure was performed. Using direct laryngoscopy or rigid bronchoscopy, the airway was evaluated in all patients. In all endotracheal intubation was performed. Four children underwent complete resection of the neck mass in the neonatal period and the neonate with haemangioma was treated conservatively with propranolol (Figs. 3-5).

Discussion

The EXIT procedure was originally reserved for management of cases with severe CDH in which tracheal clipping was introduced antenatally. During delivery, the EXIT procedure and placental support provide the surgeon with additional time for removal of the clips, bronchoscopy, intubation and surfactant administration. In the cases presented, the FE-TENDO technique with use of the PLUG balloon (PLUG – Plug the Lung Until It Grows) was used. It was introduced into the trachea of the foetus with CDH with extreme pulmonary hypoplasia in order to increase the airway pressure resulting in an increased volume of the lung and alveoli.

CDH occurs in 1:2500 to 1:5000 live births. It has a mortality rate of 20% and the degree of associated pulmonary hypoplasia and severity of pulmonary hypertension still remain a major determinant of survival. The two factors which clearly influence postnatal mortality are the timing of the termination of gestation and the presence of additional anomalies, which can increase the mortality rate up to 90%. Foetal surgery, because of its complexity and need for special instrumentation, is available only in a few centres that receive a sufficient number of cases. The optimal solution for cases with CDH is foetal surgery and delivery with the EXIT procedure at the same centre. As emergency delivery and the EXIT procedure were necessary too early (at 32 weeks of gestation) because of heart rate deceleration and CDH led to severe hypoplasia of both lungs, the probability of survival in the first case was poor.

In the second case presented, the problem of proper qualification to the EXIT procedure occurred. In the foetus, prenatally CHAOS (congenital high airway obstruction syndrome) was diagnosed which is a syndrome with near complete or complete intrinsic obstruction of the foetal airway. It is most commonly caused by laryngeal atresia, subglottic stenosis, laryngeal cyst, or laryngeal web. It may be associated with oesophageal atresia, cardiac anomalies, genitourinary anomalies, vertebral anomalies, imperforate anus, syndactyly and anophtalmia. Although the laryngeal and tracheal agenesis was not associated with other anomalies, an increase in polyhydramnios
during maternal monitoring was found which worsened the prognostic criteria and therefore the foetus did not have a chance to survive. Although all large neck masses can cause airway compression, the most common indication for the EXIT procedure is cervical teratoma (2 of 5 cases presented), lymphatic malformation (2 of the presented cases), thyroid goiter, neuroblastoma, neural tube defects and rarely haemangioma (1 of the 5 cases presented)\textsuperscript{10}. Lymphatic malformations account for only about 5% of benign tumours in infants and children. About two-thirds of reported cases are found in the head and neck, usually before the age of 2 years, and some occasionally extend to the mediastinum\textsuperscript{11,12}. Cervical teratomas are uncommon neoplasms, representing 3\% of teratomas in childhood; 5\% of cases are localized in the head and neck region with a mortality of 40-50\%. Although these lesions are histologically benign, they may be large and may cause airway obstruction (20\% of cases). Prenatal ultrasound diagnosis is possible in early pregnancy (15-16 weeks)\textsuperscript{13}. According to Laje\textsuperscript{13} both of these neck masses tend to compress the airway, but in different ways. Cervical teratomas can pull the trachea and lungs superiorly against the thoracic inlet, which can result in pulmonary hypoplasia while lymphatic malformations will not because they are softer. Cervical teratomas, in contrast to lymphatic malformations, can be invasive and destructive causing, for example, mandibular hypoplasia\textsuperscript{13}.

At the level of prenatal evaluation, the early involvement of a multidisciplinary team including neonatologist, obstetrician, anaesthesiologist, paediatric surgeon, paediatric otolaryngologist, cardiologist and radiologist is essential to correctly qualify the foetus for the EXIT procedure. Foetal ultrasonography allows visualisation of head and neck malformations and determines their size and localisation. MRI provides information about the relationship of the foetal tumour with adjacent structures such as trachea or oesophagus. It also helps to estimate the severity of the foetal airway obstruction and reveals other anomalies\textsuperscript{14,15}. Maternal monitoring for complications such as polyhydramnios and preterm labour are important in planning and desirability of the EXIT procedure. Finally, the EXIT procedure allows the paediatric otolaryngologist to provide the patency to the newborn’s airways during delivery.

**References**


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