Preoperative management of children with esophageal atresia: current perspectives

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Abstract: Esophageal atresia remains one of the most challenging congenital anomalies of the newborn. In recent years, because of the advances in prenatal diagnosis, neonatal critical care, and surgical procedures, overall outcomes have improved substantially, including for premature children. Nowadays, most of the research is focused on medium- and long-term morbidity, with particular reference to respiratory and gastroesophageal problems; the high frequency of late sequelae in esophageal atresia warrants regular and multidisciplinary checkups throughout adulthood. Surprisingly, there are few studies on the impact of prenatal diagnosis and there is continuing debate over the prenatal and preoperative management of these complex patients. In this review, we analyze the literature surrounding current knowledge on the management of newborns affected by esophageal atresia, focusing on prenatal management and preoperative assessment.

Keywords: prenatal diagnosis, esophageal atresia, tracheoesophageal fistula, ultrasound scan, tracheobronchoscopy

Introduction

With an incidence of 2.43 cases per 10,000 births, esophageal atresia (EA), with or without tracheoesophageal fistula (TEF), remains one of the most challenging congenital anomalies of newborn babies.1,2 Due to advances in prenatal diagnosis, neonatal critical care, and surgical procedures, overall outcomes have improved substantially in recent years, including for premature children.2,4 Nevertheless, EA is still associated with a life-long risk of complications, even if mortality is currently limited to those cases with associated severe life-threatening anomalies. Many unanswered questions still remain in surgical and clinical arenas. In the latter scenario, a continuing debate over the prenatal and preoperative management of these complex patients is ongoing, and more evidence-based recommendations are impelling.3–5 In this review, we analyze the current knowledge on the management of newborns affected by EA, focusing on prenatal and preoperative assessment.

Classification

The variants of EA have been described using many anatomical classification systems.1,2 To avoid ambiguity, the clinician should use a narrative description. Nevertheless, Gross of Boston described the classification system that is most often cited.1 According to Gross’ classification, the primary types of congenital EA are EA with distal TEF (85%, Gross C), isolated EA without TEF (8%, Gross A), TEF without...
atresia or H-type TEF (4%, Gross E), EA with proximal TEF (3%, Gross B), and EA with proximal and distal TEF (<1%, Gross D)² (Figure 1).

### Antenatal considerations

#### Ultrasonographic prenatal diagnosis

Ultrasonographic prenatal diagnosis of EA remains difficult and challenging, with prenatal detection rates reported as varying from 10% to 50% of cases ⁵,⁶ Multicenter studies on prenatal diagnosis are lacking, and only few national registers are currently available (Table 1).⁷⁻⁹ Traditionally, a small or absent gastric bubble in combination with maternal polyhydramnios have been the features advocated as the most sensitive sign of EA.³ Nevertheless, the combination of these two signs gave only modest predictive values, ranging from 44% to 56%.⁴⁻⁶,¹⁰ Moreover, in cases of TEF, the passage of amniotic fluid through the fistula fills the stomach and usually prevents the development of polyhydramnios. For this reason, in the study by Spaggiari et al, polyhydramnios and small/absent stomach bubbles were observed in 100% of cases of type A EA (pure EA without TEF) and only in 46.3% of cases with TEF (types B, C, D, and E, according to Gross' classification).¹¹ Similar results were reported by the 2013 French National Register, in which the prenatal suspicion of type A EA was 86% and only 12% in cases with TEF.¹¹ Considering that EA with TEF is by far the most common type of EA, this is why the overall prenatal diagnosis of EA remains quite poor.¹² The sensitivity of prenatal ultrasound (US) was, therefore, significantly higher for those scans performed in tertiary center hospitals compared with those performed in district general hospitals.¹¹⁻¹⁴ The combination of a small stomach together with the so-called pouch sign has been reported to be diagnostic for EA in few studies.¹³⁻¹⁶ This direct finding consists of the blind-ending pouch in the fetal neck or mediastinum during fetal swallowing, and it gave excellent predictive values.¹⁴ Nevertheless, the pouch sign is clearly detectable only after the 26th week of gestation (Table 2); as hypothesized by Kalache et al, in the first and early second trimester, the fetus is not able to generate sufficient pressure on swallowing to permit dilatation of the blind-ending esophagus.¹⁴ The localization of the upper pouch may also be significant in predicting the outcomes of the affected fetuses: as reported by Has et al, when the blind esophagus is localized at neck level (“upper pouch sign” or

![Figure 1](image-url)  
**Figure 1** Anatomical classification of esophageal atresia according to Gross.  
**Notes:** Anatomical classification of EA according to Gross. The primary types of congenital EA are EA with distal TEF (85%, Gross C), isolated EA without TEF (8%, Gross A), TEF without atresia or H-type TEF (4%, Gross E), EA with proximal TEF (3%, Gross B), and EA with proximal and distal TEF (<1%, Gross D).  
**Abbreviations:** EA, esophageal atresia; TEF, tracheoesophageal fistula.

<table>
<thead>
<tr>
<th>Author</th>
<th>Setting</th>
<th>Period</th>
<th>Number of EA patients</th>
<th>Prenatal detection rate (%)</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pini Prato et al⁷</td>
<td>Italian National Register</td>
<td>2012</td>
<td>146</td>
<td>29.6</td>
<td>Prenatal diagnosis significantly more frequent in Gross’ types A and B EA, compared with types C and D EA (8/10=80% vs 34/131=26%)</td>
</tr>
<tr>
<td>Leoncini et al⁸</td>
<td>Western Australian Register of Developmental Anomalies (WARDA)</td>
<td>1980⁻2009</td>
<td>260</td>
<td>7.7</td>
<td>Prenatal detection rate increased during the study period, with a corresponding decrease in the proportion of cases diagnosed in the first week of life</td>
</tr>
<tr>
<td>Garabedian et al⁹</td>
<td>French National Register</td>
<td>2008⁻2010</td>
<td>408</td>
<td>18.1</td>
<td>The morbidity rate for infants with prenatal diagnosis is significantly higher than for infants with postnatal diagnosis, with no difference in mortality rate</td>
</tr>
</tbody>
</table>

**Abbreviation:** EA, esophageal atresia.
Table 2 Prenatal findings suspected for EA

<table>
<thead>
<tr>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ultrasound</strong></td>
<td></td>
</tr>
<tr>
<td>Maternal polyhydramnios</td>
<td>High sensitivity</td>
</tr>
<tr>
<td>Absent or small gastric bubble</td>
<td>Low specificity: reported in up to 10% of pregnancies After 26th week of gestation</td>
</tr>
<tr>
<td>“Upper pouch sign” and “Lower pouch sign”</td>
<td>High specificity, but not pathognomonic for EA Predictive value for gap between esophageal pouches After 26th week of gestation</td>
</tr>
<tr>
<td>Regurgitation</td>
<td>High specificity</td>
</tr>
<tr>
<td>“Tracheal print sign”</td>
<td>The length of the tracheal widening could estimate the gap between esophageal pouches After 26th week of gestation</td>
</tr>
</tbody>
</table>

**Magnetic resonance imaging**

<table>
<thead>
<tr>
<th>Findings</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonvisualization of the intrathoracic esophagus</td>
<td>High specificity</td>
</tr>
<tr>
<td>Absent or small gastric bubble</td>
<td>More specific for pure EA (type I)</td>
</tr>
<tr>
<td>“Pouch sign” and distension of the lower esophagus</td>
<td>When visualized, distension of the lower esophagus indicates the possible presence of a tracheoesophageal fistula, while its absence favors type I EA</td>
</tr>
<tr>
<td>Tracheoesophageal fistula</td>
<td>High specificity</td>
</tr>
<tr>
<td>Tracheal bowing</td>
<td>High specificity</td>
</tr>
</tbody>
</table>

**Abbreviation:** EA, esophageal atresia.

“neck sign”), EA is more likely to be associated with long-gap atresia, whereas a mediastinal pouch (“lower pouch sign”) is more likely to be associated with distal TEF and short gap atresia. Nevertheless, Solt et al reported a small series of fetuses with polyhydramnios and a pouch sign, in which the neonatal workup ruled out EA in all cases. Eventually, Quarello et al reported an indirect sign of EA consisting of a weakness of the posterior wall of the trachea next to the esophageal defect (“tracheal print sign”); in all the six patients in their study, the length of the tracheal widening provides an estimation of the gap between esophageal pouches.

**Biochemistry of amniotic fluid**

Only a few studies have evaluated the biochemistry of the amniotic fluid in fetuses with EA. Morin et al first reported that amniotic fluid from pregnancies with fetuses with intestinal obstruction presented diminished microvillar enzyme activities, as compared to normal fetuses. In subsequent years, abnormal levels of gamma-glutamyl transpeptidase (GGT) in amniotic fluid were reported in some series. In 2011, Czerkiewicz et al proposed the EA index, corresponding to the multiplication of GGT and alpha-fetoprotein, both expressed in multiples of median; by using a threshold of EA index >3, a sensitivity of 98% and a specificity of 100% were found in the diagnosis of EA. Less encouraging results were reported by Garabedian et al, who with the same EA index observed 90% of sensitivity and 60% of specificity. The evidence with regard to routine amniotic fluid analysis in fetuses with EA is poor, including due to the fact that it has only been assessed in cases of polyhydramnios.

**The impact of prenatal diagnosis**

There has been little study of the impact of prenatal diagnosis on the prognosis of newborns with EA. Garabedian et al, analyzing data from the French National Register for infants with EA born from 2008 to 2010, concluded that patients with prenatal diagnosis have a higher morbidity rate related to the EA type (types A, B, and/or long gap), even if no difference in mortality was found. Nevertheless, prenatal diagnosis enables the research of associated abnormalities.
that could affect morbidity. Stringer et al demonstrated a high incidence of other major anomalies, particularly trisomy 18, in fetuses with suspected EA, and they indicated much poorer outcomes and a higher mortality in these patients, as compared to children with postnatal diagnosis of EA.

According to Fallon et al, when EA is suspected in a woman with polyhydramnios, prenatal management with amniotic fluid reduction should be considered to reduce the risk of preterm labor, thus exposing the infant to prematurity-related morbidity. Moreover, prenatal diagnosis allows the parents to choose for the birth to take place close to a neonatal surgery unit to avoid the problem of postnatal transfer. When an antenatal diagnosis of EA is suspected, cesarean section is recommended in the presence of associated anomalies and not to EA itself. In the Italian National Register, vaginal delivery was reported in 18 (42%) patients with prenatal suspicion of EA, with no statistically significant differences between those with and without antenatal diagnosis of EA.

**Perinatal considerations**

**Confirmation of the diagnosis**

At birth, a 10–12 French gauge nasogastric (NG) tube should be passed through the mouth into the esophagus for any infant born of a pregnancy complicated by polyhydramnios; failure to pass the NG tube beyond 11 or 12 cm has been universally recognized as the classical sign of EA. A simple X-ray of the chest and abdomen shows the tip of the catheter halted in the superior mediastinum, while gas in the stomach and intestine signifies the presence of a distal TEF. Nevertheless, in very rare instances, radiological demonstration of a catheter reaching the stomach does not exclude the diagnosis of EA, as the NG tube may take an alternative route (through the laryngeal inlet, trachea, TEF, and distal esophagus to reach the stomach), which is a rare but well-known scenario.

**Preoperative care**

The patient should be positioned sitting upright and a sump catheter should be positioned in the upper esophageal pouch to aspirate saliva continuously under low-pressure suction, in order to decrease the risk of pneumonia from aspiration of upper pouch secretions. Broad-spectrum antibiotics (ampicillin and gentamicin) and vitamin K analog are traditionally administered. Routine endotracheal intubation should be avoided because of the risk of iatrogenic gastric perforation and of increasing respiratory distress, as the abdomen becomes distended from ventilation through the TEF. In these patients, a rectal probe should be inserted to facilitate the evacuation of air and to minimize the intestinal overdistension leading to respiratory impairment.

**Associated anomalies**

A careful clinical examination should be conducted in order to rule out associated abnormalities, which are mainly responsible for the medium- and long-term prognosis in these patients. Their unequal distribution between EA patients is also important from a clinical perspective, as newborns with isolated EA without TEF (Gross A EA) exhibit anomalies in as many as 65% of cases, compared with a 10% observed in patients with TEF without atresia (Gross E EA). All patients should have an echocardiogram prior to surgery, in order to identify any structural anomaly of the heart, which is reported in up to 25% of patients. Moreover, preoperative echocardiogram could accurately detect the correct side of the aortic arch: this anomaly, although only reported in 3.6% of newborns with EA, poses a dilemma in terms of optimal surgical approach, namely the side of the thoracotomy for EA repair. When a prenatal suspicion is evident, the presence of life-threatening anomalies, including Potter’s syndrome, cerebral hypoplasia, and chromosomal anomalies (trisomy of chromosomes 13, 14, and 18), should be accurately investigated.

**Preoperative assessment**

**Contrast esophagram**

Contrast esophagram has been traditionally performed in order to detect the location of the dilated upper esophageal pouch in relation to the thoracic inlet and to detect a proximal TEF. Nevertheless, the need for contrast esophagram is still debated, as it can give false-negative results (when the fistula is occluded by mucus) or false-positive results (when the contrast identifies the tracheobronchial tree, which is more likely to be aspiration through the larynx rather than through a proximal TEF). Moreover, this procedure involves radiation hazards and may be associated with complications, including aspiration pneumonia.

**US scan, computed tomographic (CT) scan, and MRI**

Increasing evidence suggests that US scans are a useful non-invasive tool for the diagnostic assessment of newborns with EA in order to outline the tracheoesophageal anatomy. Recently, CT scans and MRIs have been proposed for newborns with EA, to identify the position of the TE fistula and anomalies of the aortic arch. Nevertheless, experience with these diagnostic tools in the preoperative assessment of newborns with EA is very limited, also for concerns regarding
neonatal transportation to the radiology department, the need for sedation, and CT-related radiation hazards. Moreover, the routine use of preoperative CT scans or MRIs in these patients is controversial, as the limited information acquired that may contribute to modifying the surgical plan can be easily obtained by preoperative tracheobronchoscopy (TBS) or intraoperatively. The only exception is represented by the extremely small subset of patients with preoperative diagnosis of right-side aortic arch and long-gap EA, in whom preoperative MRI is recommended in order to define the most appropriate surgical approach.

TBS
In recent years, preoperative TBS has received increasing attention in the evaluation of the presence of proximal TEF. However, a recent European survey demonstrated that only 43% of the respondent pediatric surgeons surveyed routinely perform preoperative TBS before EA repair. Beyond confirming the presence and location of most commonly observed lower pouch TEF, TBS enables the evaluation of vocal cord motility, to assess the presence of other specific foregut-associated anomalies (tracheomalacia, tracheal clefts, and so on), and the preoperative determination of the gap between esophageal pouches. Moreover, TBS allows the detection of rare upper pouch TEF, which could be unnoticed during surgery. Therefore, today, endoscopic assessment is an essential part of surgical planning.

Assessment of the gap
Accurate preoperative determination of the gap between esophageal pouches is mandatory in order to define the most appropriate surgical plan. Gap measurement has been largely reported in detail and should be performed at the time of preoperative TBS. When a distal TEF is evident, a rigid catheter is inserted into the upper esophageal pouch, while the tip of the bronchoscope is positioned at the level of tracheal opening of distal fistula. Chest fluoroscopy demonstrates the distance between them, which is highly representative of the gap length. Bagolan et al reported that a standardized and reproducible preoperative protocol of gap measurement enhanced the possibility of preserving the native esophagus in cases of long-gap EA.

Future perspectives on prenatal and preoperative assessment of patients with EA
EA remains one of the most challenging disorders in newborns and, although improvements have been made in the diagnosis and treatment of these patients over the years, much remains to be understood. Nowadays, research is focused on medium- and long-term morbidity, with particular reference to respiratory and gastroesophageal problems; the high frequency of late sequelae in EA warrants regular and multidisciplinary checkups throughout adulthood. Surprisingly, there are few studies on the impact of prenatal diagnosis. An accurate prenatal diagnosis is necessary in order to offer an early multidisciplinary prenatal counseling and to allow parents to choose for the birth to take place close to a high-volume tertiary-level center to offer the best of care and to avoid the problem of postnatal transfer. Unfortunately, despite recent advancements in prenatal diagnosis, the suspicion of EA is stably reported in <50% of cases of EA, with higher rate reserved only for uncommon EA variants (types A and B). Efforts should be made in order to find new signs alternative when maternal polyhydramnios is absent. Increasing evidence suggests that in case of ultrasonographic suspicion of EA, second-level MRI (including real-time MRI with cine mode) should be offered to parents. Moreover, in cases of pregnancy with polyhydramnios and suspected EA, amniotic fluid reduction should be considered to reduce the risk of preterm labor, thus exposing the infant to prematurity-related morbidity. The type of delivery (cesarean section vs vaginal delivery), according to associated anomalies, should be considered to plan appropriate perinatal care with the prompt availability of the multidisciplinary team (obstetrician, neonatologist, radiologist, neonatal surgeon, and neonatal anesthetist). At birth, an early and accurate preoperative diagnosis including TBS and, in selected case, magnetic resonance imaging is necessary to detect anatomical details such as the side of aortic arch and “long-gap” EA, in order to plan the more appropriate surgical approach before surgical repair. An early and accurate evaluation of associated anomalies, which are mainly responsible for the medium- and long-term prognosis in these patients, is also mandatory.

Author contributions
FP, ALB, SB, and DA conceptualized and designed the study, designed the data collection instruments, drafted the initial manuscript, reviewed and revised the manuscript, and approved the final manuscript as submitted. Authors declare that this manuscript has never been published and it would not be submitted to any other journal while under consideration for publication in your journal.

Disclosure
The authors report no conflicts of interest in this work.
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