Management of Tracheal Atresia with Tracheoesophageal Fistula

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Tracheal atresia is compatible with life when associated with a tracheoesophageal or bronchoesophageal fistula. Survival at birth depends on prompt diagnosis and proper airway management. The possibility of surgical correction or palliation rests on the extent of atresia present. We present a case of tracheal atresia for which a palliative operation was performed. The important features of the diagnosis and airway control of this condition are discussed.

REPORT OF A CASE

A 51-year-old woman, gravida 2, para 1, was admitted to University Hospital in premature labor with a pregnancy at 30 weeks' gestation, complicated earlier by first-trimester bleeding. Labor progressed rapidly, and a 1,250-g male infant was delivered vaginally with the aid of a pudendal block and 50 per cent nitrous oxide analgesia. Polyhydranmiosis was noticed at the time of delivery. The infant was initially cyanotic, with fair tone and reflex responsiveness, a heart rate of 100 beats/min, but little evidence of air movement despite apparently strong ventilatory efforts. The 1-minute Apgar score was 5. As the pulse rate declined to 60 beats/min, endotracheal intubation with a 3.0-mm endotracheal tube was attempted. The vocal cords were easily visualized and moved as if the infant were crying, though no sound was audible. Endotracheal intubation proved impossible, as the tube could not be passed more than a few millimeters beyond the vocal cords. Although mouth-to-tube ventilation through the partially inserted tube was attempted, the tube seemed to be completely obstructed.

The tube was removed, and ventilation via a bag and mask was begun. The heart rate increased to 120 beats/min, color improved, and ventilation was audible bilaterally on auscultation of the chest. The 5-minute Apgar score was 7.

Since ventilation via bag and mask was possible despite demonstrated subglotic obstruction, the diagnosis of tracheal web or atresia with tracheoesophageal fistula seemed likely. Also, because air was not present in the stomach despite the high airway pressures necessary for ventilation, we considered the diagnosis of esophageal atresia. Two attempts to pass an endotracheal tube blindly into the trachea via the esophagus were unsuccessful, and leaving the tube in the upper esophagus seemed only to make ventilation more difficult. Consequently, ventilation via bag and mask was continued until 50 min after birth, when an otorhinolaryngologist confirmed the presence of a tracheoesophageal fistula by placing a 5.0-mm endotracheal tube into the trachea via the esophagus under direct vision. The fistula appeared to be located approximately midway between the larynx and the carina.

The patient was admitted to the newborn intensive care unit, where continuous mechanical ventilation with PEEP of 2 cm H2O and 40 per cent oxygen was initiated. The risk of aspiration was minimized by the presence of the endotracheal tube filling the proximal esophageal segment and by frequent suctioning of the mouth and pharynx. Good respiratory status was maintained until the fourth hospital day, when oxygenation deteriorated and the murmur of a patent ductus arteriosus was heard. Congestive heart failure rapidly developed and was controlled with digoxin and furosemide. Maintenance of an adequate airway was increasingly difficult due to the instability of the endotracheal tube and the ease and frequency with which intubation of the right mainstem bronchus occurred. Therefore, the infant underwent an operation on the ninth day of life to ligate the patent ductus arteriosus and attempt definitive correction of the airway anomaly.

Anesthesia was provided with humidified nitrous oxide (2 L/min) and oxygen (1 L/min) and supplemented with d-tubocurarine. With the airway maintained by the endotracheal tube previously inserted through the tracheoesophageal fistula, the patent ductus arteriosus was ligated through a left thoracotomy. The infant’s condition improved, and laryngoscopy next showed a blind tracheal pouch a few millimeters below the normal-appearing vocal cords. Esophagoscopy revealed the esophagus to end in the upper mediastinum, with a 3-mm opening anteriorly into the trachea (Fig. 1). Maintenance of the airway during esophagoscopy was difficult, with endobronchial intubation occurring easily whenever the child was moved. At one point it was necessary to reinsert the endotracheal tube.
tube by passing a Miller #0 laryngoscope blade down the esophagus and replacing the endotracheal tube through the fistula.

Because positive-pressure ventilatory assistance would be necessary postoperatively, an attempt to provide a more secure airway was undertaken. During cervical exploration, we found that the blind tracheal pouch had no connection to the distal portion of the trachea into which the tracheoesophageal fistula opened. Tracheal reconstruction was deemed impossible due to the relatively long absent portion. Therefore, a permanent tracheal stoma was constructed by dividing the fistula and then securing the small segment of distal trachea to the skin in the sternal notch. During creation of the stoma, the airway was maintained by having a surgeon hold an endotracheal tube into place in the tracheal stump. When the stoma was complete, a shortened Shiley™ #0 tracheostomy tube was inserted and sutured in place.

This new airway proved satisfactory for continued mechanical ventilation and for subsequent general anesthesia provided for gastrostomy and correction of malrotation of the gut on the fifteenth day of life. Following this procedure, however, the infant's general condition deteriorated, with the onset of seizure activity, clotting abnormalities, and progressive congestive heart failure. The infant died on the eighteenth day of life. Postmortem examination revealed massive central nervous system hemorrhage as the cause of death.

**Discussion**

Atresia of the trachea, with or without associated esophageal defects, is an extremely rare anomaly. In a review of the literature in 1972, Hopkinson added four cases of congenital absence of the trachea to the 20 that had been recorded since 1900. Of the 24 cases he discussed, the trachea was totally absent in 19. The joined mainstem bronchi usually communicated with the esophagus, either at the level of the normal bifurcation or at a lower level (bronchoesophageal fistula); four cases with total absence of the trachea showed no communication between the respiratory tract and esophagus, a condition incompatible with life. The five cases of partial atresia all showed communication between the apex of a short length of distal trachea with the esophagus (tracheoesophageal fistula). In addition, many of these patients had other malformations of development, particularly cardiac and genitourinary. In the majority of cases the larynx was described as normal.

Although most patients with tracheal atresia have died at birth, the condition is not necessarily incompatible with life. Lyons and Bruce described an infant with an absent trachea in which ventilation was possible by bag and mask because of a communication between the esophagus and the joined mainstem bronchi. Similarly, McNie and Pryse-Davies described an infant with tracheal agenesis who survived for several hours after birth. This infant had a communication from the larynx to the esophagus and then to the mainstem bronchi.

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![Diagram of trachea and esophagus with labeled parts](http://anesthesiology.pubs.asahq.org/pdfaccess.ashx?url=/data/journals/jasa/934684/)
delayed several days for the patient presented herein, yet even with a short segment of the trachea present, considerable instability of the airway was encountered.

In selected cases of tracheal atresia surgical palliation is possible. The first surgical correction of this condition with survival beyond the immediate postoperative period was reported in 1963. This infant's trachea ended just inferior to the vocal cords; ventilation was possible because of an associated bronchoesophageal fistula. At 36 hours of life the patient underwent a cervical esophagostomy and feeding gastrostomy. The distal esophagus was divided and a respiratory passage composed of esophagus leading to the bronchi was brought out to the skin; thus, an "esophageal tracheostomy" was constructed. Other children with congenital tracheal malformations have been operated upon successfully, and a recent report details the resection of a 17-mm hypoplastic segment of trachea in a 4-month-old infant. Our patient was potentially salvagable because of the presence of distal trachea.

In summary, we wish to emphasize the following points: 1) The diagnosis of tracheal atresia with bronchoesophageal fistula or bronchoesophageal fistula should be considered when one can neither advance nor ventilate through an endotracheal tube, especially when ventilation via bag and mask proves successful; 2) initial treatment consists of assisted ventilation by bag and mask. Although attempts at endotracheal intubation should cease, a tube in the esophagus may improve ventilation; 3) an early attempt at surgical palliation (or correction) should be undertaken when the airway is inadequate or unstable. If exploration reveals the presence of a bronchoesophageal fistula, the desirability of further therapy is debatable.

REFERENCES

Unilateral Pneumothorax Following Jet Ventilation during General Anesthesia

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Jet ventilation has become an accepted and practical technique during anesthesia for laryngoscopy, fiberoptic bronchoscopy and certain head and neck surgical procedures involving invasion of the airway. However, complications such as pneumothorax and abdominal distention have been reported. This is the case report of unilateral pneumothorax secondary to endobronchial intubation with the jet catheter.

REPORT OF A CASE

A 73-year-old woman weighing 47 kg was admitted with a diagnosis of recurrent squamous cell carcinoma and scheduled for direct laryngoscopy and biopsy. She had no history of cardiopulmonary problems; preoperative roentgenogram of the chest was normal, and electrocardiogram was within normal limits. All laboratory values were within normal limits with except serum potassium, which showed slight hypokalemia (serum K+ = 3.3 mEq/l).

Hydroxyzine, 50 mg, im, and atropine, 0.4 mg, im were given 60 min before operation. Arterial blood pressure was 100/70 mm Hg. Thiopental, 160 mg, iv, was given, followed by inhalation of halothane, nitrous oxide and oxygen. The patient was then positioned with a roll under the shoulders and a Jackson laryngoscope