Olgun Sunumu – Case Report

Unexpected Subglottic Stenosis in a Newborn in Anesthesia Induction

Yenidoğanda Anestezi İndüksiyonunda Beklenmeyen Subglottik Stenoz

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Abstract

An unexpected subglottic stenosis in a neonate with multiple congenital anomalies who was scheduled for an emergency surgery of duodenal atresia was described in this case report. Duodenal atresia was diagnosed on the first day of life. After induction with sevoflurane, despite good view of the vocal cords, it was not possible to advance endotracheal tube beyond the vocal cords. After a flexible fiberoptic examination, subglottic stenosis was diagnosed and ‘an emergency tracheostomy’ decision was taken. The urgency of the operation and patent postoperative airway requirement are the key points which influence this decision.

Key Words: Subglottic stenosis, newborn, duodenal atresia

Özet


Anahtar Kelimeler: Subglottik stenoz, yenidoğan, duodenal atresi

Introduction

Subglottic stenosis (SGS) is one of the most common causes of airway obstruction in infants and children. It is the second most common cause of stridor in infants and the most common laryngotraheal anomaly requiring tracheostomy in children under 1 year of age [1]. We report the anesthetic management of an unexpected SGS in a 1 day aged newborn.

Case Report

A female neonate was born with a birth weight of 2440 g from a 29-years-old gravida 1, parity 1 mother via caesarean section at 38th week of gestation. Tetralogy of Fallot, single umbilical artery, bilateral renal-pelvic dilatation and anal atresia were diagnosed in a routine prenatal USG follow up. The family history was unremarkable with no known consanguinity. After birth, heart rate decreased to 60 min’ that was stabilized after suctioning and bag mask ventilation. The Apgar scores were 8 and 9 at 1st and 5th min, respectively. The baby was transferred to the neonatal intensive care unit because of cardiac and renal malformations. Oxygen saturation was maintained with supplemental oxygen and there has been no need for intubation. The postnatal abdominal X-ray film showed a ‘double-bubble’ sign and dilated intestinal loops which were consistent with the diagnosis of duodenal atresia. The baby was scheduled for an urgent laparotomy on the 1st day of life for correction of congenital duodenal obstruction.

Before induction of anesthesia, the baby was preoxygenated with 100% oxygen. SpO2 remained constant, over 96% and neither respiratory distress nor inspiratory stridor has been noted. Anesthesia was induced...
with sevoflurane 8% in 100% oxygen. SpO2 maintained at 99%. Then the baby was given 1 mg.kg⁻¹ fentanyl and 0.5 mg.kg⁻¹ rocuronium bromide, intravenously. Despite good view of the vocal cords, it was not possible to advance 3.0 mm, 2.5 mm and 2.0 mm internal diameter tubes beyond vocal cords because of resistance. Ear-nose and throat (ENT) consultant was invited to the operating room immediately for direct flexible fiberoptic laryngoscopy and grade-3 subglottic stenosis was diagnosed (Figure 1). The options for airway management were discussed with ENT and Pediatric surgery consultants and ‘an emergency conventional tracheostomy’ decision was found suitable because of the urgency of the operation. The otorhinolaryngologist performed a formal tracheostomy successfully and anesthesia was maintained with 50% O2-air and 2% sevoflurane under controlled ventilation.

After an uneventful sigmoid loop ostomy and duodenostomy which lasted for four hours, the baby was transferred to the neonatal intensive care unit with bag mask ventilation through the tracheostomy cannula.

**DISCUSSION**

Congenital subglottic stenosis is the third most common congenital laryngeal anomaly, after laryngomalacia and vocal fold paralysis [1]. The estimated incidence of congenital pediatric subglottic stenosis is 5% of all cases and below 1% when very low birth weight infants are excluded [1, 2]. The majority of the cases of subglottic stenosis are acquired, most commonly associated with prolonged endotracheal tube intubation, laryngopharyngeal reflux, infection and the associated inflammatory response and generally a congenital stenosis is less severe than acquired one [3].

The diameter of normal subglottic lumen is 4.5 to 5.5 mm in a full-term neonate and approximately 3.5 mm in a preterm baby. A subglottic airway diameter of 4 mm or less in a full-term infant or 3 mm or less in a premature infant is considered narrow and consistent with a diagnosis of subglottic stenosis [1]. Several staging systems have been created to describe and guide treatment options of subglottic stenosis and the Myer-Cotton system is one of the most reliable and widely used staging system [4].

Antenatal diagnosis of congenital airway obstruction syndrome is difficult. It is associated with normal larynx to the level of the vocal cords and little airway below this [5]. Failure to intubate beyond the vocal cords is frequently reported. Recurrent or persistent croup is typical, particularly in infants younger than 6 months. Although laryngeal anomalies frequently are first symptom-atic, as a result of the inflammation caused by a common respiratory tract infection, our patient did not have any obstructive symptom as suprasternal and subcostal retractions, dyspnea or tachypnea. And we did not observe any inspiratory stridor, respiratory distress or saturation problem which could warn us about subglottic stenosis.

The presence of subglottic pathology in a child can be co-incidental with genetic syndromes or non-syndromic congenital anomalies such as esophageal and duodenal atresia [6], Down syndrome [7], VATER (vertebral, anorectal, tracheo-esophageal, and renal defects) syndrome [8], Fraser syndrome [9], Keutel syndrome [10]. The presence of a subglottic pathology should be kept in mind in a child with genetic syndromes or other non-syndromic congenital anomalies, especially when these involve the mediastinum, cardiovascular or esophageal anomalies. Our patient’s associated anomalies were tetralogy of fallot, single umbilical artery, bilateral renal-pelvic dilatation, duodenal and anal atresia.

When encountered in perioperative period, tracheostomy decision is hard to take, especially in newborns with congenital subglottic stenosis. The number of complications is higher during neonatal period. This can in part be explained by using smaller tubes and associated diseases. Fewer than 50% of children require a tracheotomy and most of them can be decannulated once the airway improves sufficiently [11]. In newborns, if the surgery is not urgent, the anesthesia should be postponed and management of subglottic stenosis should be planned in elective conditions [12]. In our case, as the surgery for duodenal atresia was urgent, we could not postpone the procedure.

As described in this case report, management of an unexpected subglottic stenosis in a newborn is critical...
and the anesthetist should be aware of possible subglottic stenosis in babies with difficulties of intubation. Infants with multiple congenital anomalies may have subglottic stenosis although they have no any warning signs such as stridor or respiratory difficulty. In such situations, the urgency of the operation and postoperative safe airway requirement are the key points which influence the decision.

REFERENCES