Cystic hygroma; difficult airway and the anesthetic considerations

Nighat Abbas¹, Arif Iftikhar Mallhi², Syed Sajjad Alam³, H.M. Abid Mughal³

ABSTRACT

A 5 days old baby was referred from a local hospital for evaluation and treatment of a huge swelling on the right side of the neck extending beyond the midline in front and with difficulty by her mother to feed her since birth. It was clinically diagnosed to be cystic hygroma and a decision was made to further evaluate by doing magnetic resonance imaging under general anesthesia in order to see the extent of the mass and to plan for either sclerotherapy or surgical excision. Difficult intubation was anticipated and the baby was intubated in main operating room with inhalation induction and maintenance with sevoflurane without giving muscle relaxants. The 30 min of the procedure period was smooth without any complication. The baby was extubated in the main operating room complex.

Key words: Cystic hygroma; Difficult airway; Neonate

INTRODUCTION

Hygroma is a Greek word which means water containing tumor. Cystic hygroma is a benign tumor which is composed of large cysts and classified as capillary, cavernous and cystic lymphangiomas. It is also classified on the basis of size of the cysts as macrocystic, macrocystic and mixed lymphangiomas. A microcystic lymphangioma consists of cysts < 2 cm, macrocystic lymphangioma consist of cysts > 2 cm and mixed lymphangiomas consist of cysts of variable sizes.¹²  The airway and anesthetic management in neonates and adults has many significant differences. Maintaining the airway of neonate and young infants with large neck mass like a huge cystic hygroma can be a challenge for any anesthetist, as airway obstruction, hypventilation and hypoxemia and sudden collapse are very common risk factors due to tumor extension in the neck, airway and thorax. We describe the difficulties encountered in intubation and anesthetic management in our neonatal patient.
normal with no local rise of temperature.

Pre-anesthesia assessment was carried out with systemic review i.e. electrocardiography was within normal limits and normal S1+S2+0 cardiac sounds on auscultation, chest was clear with normal vesicular breathing and equal air entry bilaterally on auscultation, thyromental distance was less than 3 fingers, labs (complete blood count, erythrocyte sedimentation rate etc.) were normal in range and chest x-ray was done to exclude the presence of chest infection and intrathoracic extension of the tumor.

Keeping in mind the suspected difficult airway we kept ready a difficult airway cart that contained face mask of various sizes, oropharyngeal airways of assorted sizes, nasopharyngeal airways, endotracheal tubes size 2.5, 3 and 3.5, stylet, suction, bougie, short handle laryngoscope, straight blade, videolaryngoscope, laryngeal mask airway No. 1 and 1.5.

The child was premedicated with inj glycopyrrolate 0.005 mg/kg (0.014 mg). A shoulder roll was used to keep the child at optimal laryngoscopic position and preoxygenated with high flow 100% oxygen into the breathing system in order to increase oxygen reserve which might allow more time for laryngoscopy and tracheal intubation.

The child was induced with sevoflurane in oxygen by an effective face mask seal. A straight blade laryngoscope was used as child had an anterior and cephalad placed larynx and epiglottis was difficult to be visualized. At first attempt, laryngoscopy was difficult as the epiglottis and vocal cords were not visualized due to displacement of soft tissue to the left side. The child was ventilated with mask and a second attempt at laryngoscopy was made by shifting the soft tissues to the right side. It helped to visualize the glottis with assistance and the trachea was successfully intubated with uncuffed endotracheal tube (ETT) size 3 mm; fixed just 1 cm beyond the vocal cords in order to avoid accidental extubation. The child was deeply sedated with sevoflurane showing no cough reflex and did not require any muscle relaxant.

The child was transported for magnetic resonance imaging (MRI) procedure room after taking transport consent with parents of the child in an incubator with monitoring all vitals and electrocardiography with assisted ventilation. After taking the child on MRI table, all vital signs were monitored with MRI compatible monitoring system and anesthesia was maintained with isoflurane in oxygen with controlled ventilation with Jackson Rees modification of Ayre's t piece. At the end of procedure, the child was shifted back to the main operating room for extubation with all precautions taken during transportation to MRI room. The child was extubated after return of adequate muscle power, respiratory effort, good cry and limb movements.

DISCUSSION

Cystic hygroma, also called cavernous hemangioma, is caused by blockages in the lymphatic system and is a histological benign congenital tumor developed in 9th and 16th week of pregnancy.3,4

Embryologically, surrounding normal tissues are penetrated by sprouting endothelial membrane sequestered lymph vessels that forms fimbriate causing canalization and production of large multiloculated cysts filled with serous secretion.5 The condition may develop due to some genetic disorder like Turner syndrome, trisomy 13,18 or 21, Noonan syndrome or may be due to environmental factors like viral infections, exposure to drugs or alcohol passed from mother to baby during pregnancy. Approximately half of all fetuses with a cystic
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Cystic hygroma have chromosomal abnormalities therefore other related abnormalities should also be ruled out.4

Cystic hygroma usually presents in neonates and during early infancy but can develop at birth or with obstructed labor.6 The most prominent sign of cystic hygroma is presence of a soft and spongy lump which commonly appears on neck that can interfere with normal breathing and swallowing.7

Securing the airway was the prime consideration in our case. A difficult airway is a clinical situation in which a conventionally trained anesthesiologist experiences difficulty with facemask ventilation, or difficulty with tracheal intubation or both. A difficult laryngoscopy is when it is impossible to view any part of the vocal cords during direct laryngoscopy. Managing a difficult airway in newborn can be a nightmare for any physician. Intubation and managing the secure airway in newborn is difficult, and the success ratio of intubation is only 60% on first attempt. In addition, lower functional residual capacity makes them prone to rapid development of hypoxia, so it is recommended to limit the duration of laryngoscopy and intubation to less than 20 sec.8

In babies with anticipated difficult airway intubation should be carried out under general anesthesia or deep sedation, preferably maintaining spontaneous ventilation until trachea is successfully intubated.9

Extubation should be done after return of adequate muscle power, respiratory efforts, response to commands, and when there is no cardiopulmonary instability, and the cry in case of newborn or infants is good.10

CONCLUSION

Difficult airway due to head and neck mass effect can be a challenge to any anesthetist. Inhalational induction is the preferable techniques. Endobronchial intubation along with intra-operative management of intubation, accidental extubation and anticipation of possible post-operative complications should be considered. A definite plan to deal with ‘can’t intubate, can’t ventilate’ (CICV) situations, availability of appropriate equipment and critical skills are fundamental for the airway management in such cases. Similarly, during extubation, intact airway reflexes, GCS level, the respiratory effort and postoperative management should be considered to improve the prognosis of the patient.

Conflict of interest: Nil

Authors' Contribution:
NA: Supervised the case, manuscript editing & reviewing
AIM: pre-anaesthesia assessment, conduction of anesthesia
SSA: Concept, assisted the case, manuscript writing, editing
HMAM: Assisted the case

REFERENCES