REVIEW PAPER

Subglottic stenosis in paediatric patients

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ABSTRACT

Subglottic stenosis (SGS) is a complex and challenging clinical entity for paediatric otolaryngologists. Subglottic stenosis is a congenital or acquired lesion characterised by narrowing of the airway extending from the lower free border of the vocal folds to the lower border of the cricoid cartilage. The most common cause of SGS is prolonged endotracheal intubation. The treatment option for SGS is often surgery, which may be endoscopic or open. Nowadays, the endoscopic approach is reaching its limit, and open surgery is advised only in failed cases of endoscopic approach. In this review article, we discuss the current concept and recent advances in the diagnosis and treatment of SGS.

KEY WORDS: subglottic stenosis, paediatric patient, endoscopic approach, laryngotracheal surgery.

INTRODUCTION

Subglottic stenosis (SGS) is defined as narrowing of the subglottis, which is a part of the larynx extending from the lower margin of the true vocal cords to the inferior border of the cricoid cartilage [1]. Subglottic stenosis can be an acquired clinical entity occurring after prolonged intubation or a congenital condition leading to respiratory distress during the neonatal period. There has been significant advancement in the investigations and treatment of the SGS in children. The management of the SGS needs a team approach, which includes a paediatric otolaryngologist, anaesthesiologist, paediatrician, pulmonologist, and intensive care specialist. The incidence of SGS is around 1 to 2% among intubated neonates and 11% of the intubated children below five years old [2]. The rate of SGS increases by 50% for every five days of intubation in children below five years old [3]. A premature baby can tolerate prolonged intubation better than a term baby. This review article focuses on the etiopathogenesis, clinical presentations, and current management of SGS in children, with special emphasis given to the clinical experiences.

ETIOPATHOGENESIS OF SUBGLOTTIC STENOSIS

The etiopathogenesis of SGS is complex. The aetiology of congenital SGS is attributed to the developmental failure during pregnancy and is often associated with congenital head and neck lesions and syndromes like 22q11 deletion, Down syndrome, and CHARGE syndrome [4]. Acquired SGS is often due to prolonged ventilation at the intensive care unit. The subglottis of the larynx is vulnerable to intubation trauma due to its...
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The clinical manifestations of SGS are directly related to the degree of stenosis. Grade I SGS is often asymptomatic, but the symptoms are aggravated during upper airway infections with respiratory distress and even stridor. Grade III and IV SGS often present with air hunger, dyspnoea, biphasic stridor, suprasternal, intercostals, and diaphragmatic retractions. Recurrent episodes of croup may give rise to suspicion of SGS. It is often important for the clinician or paediatrician to urgently identify a compromised airway that might lead to rapid deterioration and needs prompt intervention to avoid a catastrophic situation. Extubation failure in a child often excludes the non-laryngeal causes like nasal obstruction, glossoptosis, and tracheobronchomalacia for respiratory distress. It is always advisory to document difficult intubation, history of prolonged intubation, as well as failed extubation, which give clues as to the cause of SGS. If there is any past history of airway surgery, it should always be documented with the type and timing of surgery. The clinical conditions that cause stridor are the differential diagnosis of SGS. These clinical situations are croup, subglottic haemangioma, subglottic cyst, vocal cord paralysis, complete tracheal rings, and bacterial tracheitis. Congenital SGS is the third most common congenital lesion of the larynx [7]. Subglottic stenosis is considered as congenital when there is no history of endotracheal intubation or any acquired causes of SGS. The clinical presentations of congenital SGS are directly related to the degree of narrowing of the subglottis. In severe congenital SGS, stridor is present at the time of birth, whereas in less severe, the symptoms are seen after few months of birth. Congenital SGS is less common than acquired and can only be diagnosed before its first intubation in child.

CLINICAL PRESENTATIONS

The clinical manifestations of SGS are directly related to the degree of stenosis. Grade I SGS is often asymptomatic, but the symptoms are aggravated during upper airway infections with respiratory distress and even stridor. Grade III and IV SGS often present with air hunger, dyspnoea, biphasic stridor, suprasternal, intercostals, and diaphragmatic retractions. Recurrent episodes of croup may give rise to suspicion of SGS. It is often important for the clinician or paediatrician to urgently identify a compromised airway that might lead to rapid deterioration and needs prompt intervention to avoid a catastrophic situation. Extubation failure in a child often excludes the non-laryngeal causes like nasal obstruction, glossoptosis, and tracheobronchomalacia for respiratory distress. It is always advisory to document difficult intubation, history of prolonged intubation, as well as failed extubation, which give clues as to the cause of SGS. If there is any past history of airway surgery, it should always be documented with the type and timing of surgery. The clinical conditions that cause stridor are the differential diagnosis of SGS. These clinical situations are croup, subglottic haemangioma, subglottic cyst, vocal cord paralysis, complete tracheal rings, and bacterial tracheitis. Congenital SGS is the third most common congenital lesion of the larynx [7]. Subglottic stenosis is considered as congenital when there is no history of endotracheal intubation or any acquired causes of SGS. The clinical presentations of congenital SGS are directly related to the degree of narrowing of the subglottis. In severe congenital SGS, stridor is present at the time of birth, whereas in less severe, the symptoms are seen after few months of birth. Congenital SGS is less common than acquired and can only be diagnosed before its first intubation in child.

GRADING OF SUBGLOTTIC STENOSIS

The classical grading of SGS (Fig. 1) was described by Cotton and approved worldwide. The modified Cotton Meyer grading (Fig. 2) based on the percentage of obstruction calculated on the basis of passing an endotracheal tube through the stenosis, resulting in the approximate diameter of stenotic segment divided by the age-appropriate endotracheal size [8]. There are some disadvantages of this grading system. Vocal cord status has an impact on the treatment of SGS. The vocal cord immobility with grade I stenosis will not provide an adequate airway. Other factors like length of SGS have an impact on the treatment. A severe SGS with a thin scar web is usually treated easily by endoscopic method, whereas a long but a less narrow segment is often resistant to endoscopic ap-

<table>
<thead>
<tr>
<th>Classification</th>
<th>From</th>
<th>To</th>
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<tbody>
<tr>
<td>Grade I</td>
<td>No obstruction</td>
<td>50% obstruction</td>
</tr>
<tr>
<td>Grade II</td>
<td>51% obstruction</td>
<td>70% obstruction</td>
</tr>
<tr>
<td>Grade III</td>
<td>71% obstruction</td>
<td>99% obstruction</td>
</tr>
<tr>
<td>Grade IV</td>
<td>No detectable lumen</td>
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FIGURE 1. Cotton Meyer grading of SGS
proach. The consistency of SGS, e.g. granulation tissue or scar tissue, also has an impact on the treatment.

DIAGNOSIS

The diagnosis of SGS (Fig. 3) is done by proper history taking and physical examination of the child. History should be taken regarding birth injury, prolonged intubation, prematurity, and other congenital anomalies, and physiologic indicators of respiratory or cardiovascular status should be documented. The duration and onset of stridor should be documented. The type of stridor, e.g. inspiratory, expiratory, and biphasic, should also be documented. The voice quality of the child is also important to give an idea of the location of the pathology [9]. Feeding problems should be documented. History of extubation failure or non-laryngeal causes like nasal obstruction, glossoptosis, and tracheobronchomalacia should also be documented. History of difficult intubation or failed extubations supports laryngotracheal stenosis. The situation, like the current ventilator requirement of the child, determines whether early reconstruction is needed or not. Any history of endoscopic or open airway surgery should be documented, including timing and reasons for failure of the surgery. In the case of tracheostomised children (Fig. 4), the size of the tracheostomy tube, tolerance to the speech valve, and the possibility of tracheostomy tube plugging should be documented. The clinical conditions causing stridor should be considered for differential diagnosis, like croup, subglottic haemangioma, subglottic cyst, bacterial tracheitis, vocal cord paralysis, and complete tracheal rings.

IMAGING

The imaging evaluation provides detailed information regarding the site and length of the stenotic segment at

<table>
<thead>
<tr>
<th>Patient Age</th>
<th>Percentage of Obstruction Actual Endotracheal Tube Size:</th>
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<tbody>
<tr>
<td></td>
<td>ID = 2.0</td>
</tr>
<tr>
<td>Premature</td>
<td>No obstructed</td>
</tr>
<tr>
<td>0-3 mo</td>
<td>58</td>
</tr>
<tr>
<td>3-9 mo</td>
<td>68</td>
</tr>
<tr>
<td>9 mo – 2 yr</td>
<td>75</td>
</tr>
<tr>
<td>2 yr</td>
<td>80</td>
</tr>
<tr>
<td>4 yr</td>
<td>84</td>
</tr>
<tr>
<td>6 yr</td>
<td>86</td>
</tr>
</tbody>
</table>

FIGURE 2. Modified Cotton Meyer grading of SGS

FIGURE 3. Endoscopic picture showing SGS
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The diagnosis of the SGS is usually confirmed by direct laryngoscopy. A routine computed tomography (CT) scan or magnetic resonance imaging (MRI) is advised for most children with SGS. Magnetic resonance angiography (MRA) is usually done when suspected for vascular anomaly or tumour causing airway obstruction. An MRI or CT scan is done when direct laryngoscope reveals near complete or complete obstruction at the subglottis, to assess the length of the stenotic segment [10].

ENDOSCOPIC EVALUATION

The gold standard evaluation for laryngotracheal airway stenosis is endoscopic assessment. Rigid bronchoscopy and microlaryngoscopy will identify the site and degree of stenosis from larynx to carina. If a patient has a tracheostomy tube, attention should be paid to the suprastomal area to rule out any suprastomal collapse, suprastomal granuloma, and any skin tract. The distal and middle tracheal assessment rules out any complete tracheal rings, a tracheoesophageal fistula, vascular compression, or tracheomalacia. The carina and main bronchi should be examined.

TREATMENT

The treatment on SGS is based on history, severity of the clinical presentations (breathing difficulty), age of the child, vocal cord mobility, previous intervention, co-morbidities like developmental, neurological, and aspiration, and other associated airway lesions. Taking the above considerations along with direct laryngoscopy findings and an imaging report, the appropriate decision is taken for treatment of SGS. A child presenting with respiratory distress in the emergency room require urgent attention and intervention. A sign of impending disaster is increased effort of breathing despite normal oxygen saturation. In this situation, the child may suddenly decompensate. Here, a quick decision is needed for secure airway. The safest area in which to provide a secured airway is the operating room, where direct laryngoscopy or rigid bronchoscopy can be done in the case of difficult intubation. A tracheostomy set should be ready for emergent tracheostomy if the above procedures are unsuccessful. Because the airway of the child is collapsible, cricothyroidotomy is unsafe and is not a suitable procedure, so it should be avoided in children. Inhalation nebulisation with adrenaline and use of heliox may help to reduce breathing difficulties before intervention to provide a secure airway. Children with SGS with grade I or mild grade II without tracheostomy may not need surgery and can be observed [11]. In children with very mild stridor, never hospitalised for imminent airway compromise. As children have reactive larynx, watchful waiting and avoiding surgery those have tracheostomy. A larynx is referred to as “reactive” when it is associated with inflammation and granulations. Paediatric patients tend to heal poorly after surgery, so open airway reconstructive surgery should be avoided until the larynx is no longer “reactive”. Laryngotracheal reconstructive surgery is contraindicated in: children with low weight, e.g. less than 10 kg, recurrent aspiration pneumonia, children with neurological lesions with high potential for recurrent aspiration, and those with severe pulmonary disease [11].

MEDICAL TREATMENT

Medical treatment is useful in a few selected cases. Intravenous or inhaled corticosteroids are indicated if SGS is diagnosed early. It influences the inflammatory process at the site of the lesion. Beclomethasone aerosol every six hours improves the patient’s symptoms. Intravenous hydrocortisone prevents suprastomal granulations [12].

SURGICAL TREATMENT

The two surgical options in SGS are: endoscopic or open. Endoscopic surgery is usually helpful in the treatment of grade I or II. Grade III or IV SGS need open surgical treatment like laryngotracheal reconstructions (LTR), laryngotracheoplasty (LTP), or cricotracheal resection (CTR). But nowadays, with the advancement of technology, the endoscopic approach has reached its limit, and open surgery is only done in case of failed cases of endoscopic approach.

MICRODEBRIDER

The microdebrider is usually used in endoscopic sinus surgery but may also be suitable for laryngeal endoscopic sinus surgery by using skimmer blades [13]. The microdebrider is used in cases of laryngeal papillomatosis, and it can be used in subglottic granuloma or fibroma, haemangioma, and subglottic cysts. The advantage of the microdebrider is its accuracy of tissue excision without damaging adjacent tissue.
**BALLOON DILATION**

Balloon dilation is an effective endoscopic surgical option and it is an alternative to open reconstructive surgery. It has a 66% success rate in preventing tracheostomy or LTR [14]. It has also a relatively low complication rate. Serial balloon dilation is a safe and effective endoscopic method for treating SGS.

**BOUGIE DILATION**

Subglottic stenosis is treated with serial dilation by bougie (steel dilator). Its use is declining at present, and most surgeons prefer endoscopic balloon dilations. However, the results of bougie dilation are similar to the balloon dilation.

**TRACHEOSTOMY**

Tracheostomy (Fig. 3) is sometimes the safest way to serve to airway in SGS. It may serve as a bridge before doing laryngotracheal surgery. Tracheostomy is important particularly when the larynx appears inflamed and active leading to failure of the surgery.

**OPEN SURGERY**

If the endoscopic approach fails or the appropriate success rate is not achieved, wide arrays of open airway reconstructive methods are performed to treat the laryngotracheal stenosis. Open airway laryngotracheal reconstruction can be done either as a single- or double-stage procedure. The single-stage procedure is not recommended in children with a history of difficult intubation, poor pulmonary function, reconstruction failure, or sedation problems [15]. In the double-stage procedure, first the patient requires tracheostomy, and the decannulation is done in a second stage.

**LARYNGOTRACHEAL RECONSTRUCTION**

Children with severe SGS benefit maximally from LTR by using anterior costal cartilage grafting, and it is also useful in failed cases of endoscopic management. Its success rate is more than 90% [16]. It can be a single- or double-stage procedure. The most common graft material used in LTR is costal cartilage, which gives excellent long-term outcomes [17]. The costal cartilage is readily available, easily craved, and easy to make into a flange so that it decreases the risk of graft prolapse into the laryngotracheal airway [17]. Sometimes the thyroid ala is harvested easily from the same surgical field. It has the limitation of not being amenable to being craved with a flange. It is only harvested when a small graft is needed. Conchal cartilage of the pinna is also used for LTR, with successful outcome in one study [18].

**ANTEOR CRICOID SPLIT**

It is an alternative procedure to tracheostomy in premature infants with prolonged intubation. It was initially started as an open surgical procedure to decompress the pressure of the endotracheal tube on the subglottic oedema. Presently, endoscopic anterior cricoid splits along with balloon dilations are reported with 83% success rate [19]. Larger studies are needed to establish this novel technique.

**CRICOTRACHEAL RESECTION**

The aim of this surgical procedure is to remove the stenotic segment of the airway, and the reunion of the healthy superior and inferior segments [20]. Cricotracheal resection is indicated in severe SGS or structurally inadequate subglottis and in patients who have undergone previous airway reconstructive surgery. The relative contraindications to CTR are low-grade SGS, stenosis within 3 mm of the vocal cords, or situations that impair mobilisation of the trachea, such as past history of distal tracheal surgery or injury to the tracheoosophageal septum [15]. Cricotracheal resection is a more technically challenging surgery in comparison to LTR, and there is disruption of the laryngeal framework and its supportive parts, no donor site morbidity, and decannulation may be successful. The disadvantages of CTR are the risk of catastrophic dehiscence of the repair and the chance of injury to the recurrent laryngeal nerve.

**SLIDE TRACHEOPLASTY**

Slide tracheoplasty done by extending through the anterior cricoid and effectively treating the patient with SGS. This approach may be cervical or transthoracic [21].

**STENTING**

Stenting with a Montgomery T-tube is relatively safe in children but is associated with aspiration in 11.5% and granulation in 16% of the cases [22]. The use of a silicone laryngotracheal mould reduces the formation of granulation, and it can stay for several months in difficult cases of SGS. The only prerequisite for this procedure is a tracheostoma. If excessive granulation tissue is seen on the cartilage graft after removal of the laryngotracheal mould (LT mould), mitomycin-C may be applied after removal of granulations and the mould can be reinserted for 1-2 months and an effective method for decannulation [23].

**POSTOPERATIVE COMPLICATIONS**

The complications after airway surgery may occur during surgery or after a procedure that includes desatu-
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RATION, bleeding, wound infections, dehiscence, and graft dislodgment. The single-stage reconstruction of airways leads to additional complications like prolonged sedation/paralysis, unplanned extubation, oedema in the airway, and severe narcotic withdrawal.

CONCLUSIONS

Subglottic stenosis is a challenging clinical entity. Inadequate airway due to SGS in children is a life-threatening situation. It is often good to avoid tracheostomy in children. After successful surgical treatment, most of the children are decannulated. An experienced team and advanced technology will be safest for obtaining optimum results.

DISCLOSURE

The authors declare no conflict of interest.

REFERENCES