Feasibility of Surgical Treatment for Laryngomalacia Using the Spontaneous Respiration Technique

Hyunkyung Cha1,* · Doh Young Lee2,* · Eun-Hee Kim3 · Ji-Hyun Lee3 · Young-Eun Jang3 · Hee-Soo Kim3 · Seong Keun Kwon1

1Department of Otorhinolaryngology-Head and Neck Surgery, Seoul National University Hospital, Seoul National University College of Medicine, Seoul; 2Department of Otorhinolaryngology-Head and Neck Surgery, SMG-SNU Boramae Medical Center, Seoul National University College of Medicine, Seoul; 3Department of Anesthesiology and Pain Medicine, Seoul National University Hospital, Seoul National University College of Medicine, Seoul, Korea

Objectives. In this study, we review our institutional experience with pediatric laryngomalacia (LM) and report our experiences of patients undergoing supraglottoplasty using the spontaneous respiration using intravenous anesthesia and high-flow nasal oxygen (STRIVE Hi) technique.

Methods. The medical records of 29 children with LM who visited Seoul National University Hospital between January 2017 and March 2019 were retrospectively reviewed. Surgical management was performed using the STRIVE Hi technique. Intraoperative findings and postoperative surgical outcomes, including complications and changes in symptoms and weight, were analyzed.

Results. Of the total study population of 29 subjects, 20 (68.9%) were female. The patients were divided according to the Olney classification as follows: type I (n=13, 44.8%), II (n=10, 34.5%), and III (n=6, 20.7%). Twenty-five patients (86.2%) had comorbidities. Seventeen patients (58.6%) underwent microlaryngobronchoscopy under STRIVE Hi anesthesia. Four patients with several desaturation events required rescue oxygenation by intermittent intubation and mask bagging during the STRIVE Hi technique. However, the procedure was completed in all patients without any severe adverse effects. Overall, 15 children (51.7%) underwent supraglottoplasty, of whom 14 (93.3%) showed symptom improvement, and their postoperative weight percentile significantly increased (P=0.026). One patient required tracheostomy immediately after supraglottoplasty due to associated neurological disease.

Conclusion. The STRIVE Hi technique is feasible for supraglottoplasty in LM patients, while type III LM patients with micrognathia or glossoptosis may have a higher risk of requiring rescue oxygenation during the STRIVE Hi technique.

Keywords. Laryngomalacia; Airway Management; STRIVE Hi; Anesthesia; Surgical Procedure; Supraglottoplasty

INTRODUCTION

Laryngomalacia (LM) is the most common (45%–75%) cause of congenital stridor, characterized by inward collapse of laryngeal structures, resulting in a narrow air passage and turbulent airflow that causes inspiratory stridor [1]. It generally worsens with feeding, crying, placing in the supine position, and agitation, and may lead to feeding problems, failure to thrive, and significant respiratory distress [2]. Olney et al. [1] categorized LM into three types according to the site of obstruction and the surgical procedure most likely to affect the outcome: type I, inward collapse of the aryepiglottic fold during inspiration; type II, long tubular epiglottis with short aryepiglottic folds; and type III, overhanging epiglottis that collapses posteriorly [3]. Diseases associated with LM include gastroesophageal reflux disease (65%–100%), congenital anomalies/syndromes (8%–20%), neu-
rological diseases (20%-45%), cardiac diseases (10%), and synchronous airway lesions (SALs; 7.5%-64%) [2]. Although most LM patients are managed with conservative medical therapy, patients with severe disease whose symptoms worsen despite treatment require surgical intervention under general anesthesia [4]. The conventional method using closed-circuit anesthesia with an endotracheal tube has several constraints for pediatric patients. Due to the small surgical field and anatomy of the pediatric airway and the thickness of the tube, endotracheal intubation results in the surgical field being covered by the tube, making it difficult to observe the surgical site and SAL in the lower airway [5]. For a wider surgical view, the intermittent apneic technique with jet ventilation was introduced for tubeless airway surgery. However, this technique requires repeated intubation, which is associated with risks of laryngeal injury, hypoxemia, and hypercapnia, and is a time-consuming procedure [5,6]. In addition, jet ventilation is associated with a risk of barotrauma due to high airway pressure [5,7,8]. Evaluating dynamic changes in the larynx of patients with LM is limited if a muscle relaxant is administered when performing anesthesia.

The spontaneous respiration using intravenous anesthesia and high-flow nasal oxygen (STRIVE Hi) technique is a tubeless, spontaneous-respiration anesthesia technique that uses total intravenous anesthesia in combination with high-flow nasal oxygen [8]. Using the tubeless self-respiration technique, we were able to evaluate dynamic obstruction of the upper airway in our patients, leading to an accurate diagnosis before performing supraglottoplasty. High-flow nasal oxygen has benefits with regard to prolonged apnea and reduction of the number of intermittent intubations [9]. The results of the procedure could be evaluated intraoperatively and further adjustments could be performed as necessary. A final dynamic evaluation of the surgical results was also possible using STRIVE Hi. This study investigated the characteristics of patients with LM who were referred to our tertiary hospital, and assessed the safety and efficacy of the STRIVE Hi technique and surgical outcomes of supraglottoplasty using this technique.

**Highlights**

- We analyzed the feasibility of the spontaneous respiration using intravenous anesthesia and high-flow nasal oxygen (STRIVE Hi) technique for the surgical management of laryngomalacia (LM).
- A relatively high proportion of LM patients in this study had multiple comorbidities.
- Approximately 73% of patients (13/17) did not need rescue oxygenation during the STRIVE Hi technique, and most of them (93.3%) showed symptom improvement without any complications.
- Type III LM patients with micrognathia or glossoptosis may have a higher risk of requiring rescue oxygenation during the STRIVE Hi technique.

**MATERIALS AND METHODS**

**Subjects**

A retrospective review of medical records was performed for children with LM diagnosed and treated at Seoul National University Hospital from January 2017 to March 2019. Each subject underwent an evaluation that included history-taking, physical examination, and flexible laryngoscopy. Patients who were not followed up were excluded. The study population consisted of 29 patients diagnosed with LM by flexible laryngoscope examination. Type classification was performed based on the transnasal fiberoptic exam according to Olney’s classification [1]. The study protocol was approved by the institutional review board of the Clinical Research Institute at Seoul National University Hospital (IRB No. 2004-226-1119). All methods were performed in accordance with the approved guidelines and the Declaration of Helsinki. All personal information was kept confidential as required.

**Surgical techniques**

Microlaryngobronchoscopy was considered for airway evaluation and accurate diagnosis in patients whose symptoms could not be controlled by conservative medical therapy. After conducting microlaryngobronchoscopy, the surgery, including supraglottoplasty, was determined as necessary. Indications for supraglottoplasty included severe airway obstruction presenting with significant chest retraction, bodyweight percentile <5%, or failure to thrive [10]. Surgery was performed by a single surgeon (SKK). Microlaryngobronchoscopy and supraglottoplasty were performed using the STRIVE Hi anesthesia technique. The main surgical technique used in this study for type I cases was making an incision on the short aryepiglottic fold and excising the redundant aryepiglottic fold (Fig. 1A). The posterior commissure mucosa was left untouched to avoid stenosis. For type II patients, the lateral side of the tubular epiglottis was vaporized in addition to the procedure for type I LM (Fig. 1B). For type III patients, epiglottopexy was performed using a CO2 laser to burn the mucosa of the tongue base and lingual surface of the epiglottis, and a suture was placed in between (Fig. 1C).

**STRIVE Hi anesthetic technique**

Starting with 3 minutes of preoxygenation, an AIRVO humidifier high-flow system (Fisher & Paykel, Auckland, New Zealand) was used. The flow rate was set to 2 L/min/kg and the FiO2 was set to 1.0 and routinely reduced to <0.3 while using the CO2 laser. Intravenous (IV) 1% lidocaine and 1% propofol were administered during induction, and general anesthesia was maintained by IV infusion of propofol (100–250 µg) and remifentanil (0.01–0.1 µg/kg/min). The sedative infusion rate was maintained at a level to avoid undesirable events (patient movement, laryngotraheal spasm) while preserving the patient’s self-respiration. Noninvasive blood pressure and pulse oximetry (SpO2), as well
Fig. 1. Supraglottoplasty techniques for each type of laryngomalacia. (A) Type I, excision of short and redundant aryepiglottic fold. (B) Type II, lateral partial epiglottectomy in addition to the procedure used for type I. (C) Type III, excision of short aryepiglottic fold with epiglottopexy of after laser vaporization of epiglottis.
as electrocardiography and oxygen reserve index (ORI) were monitored. A transcutaneous carbon dioxide (TcCO₂) monitor was attached to the right side of the chest instead of end-tidal carbon dioxide monitoring [5].

Outcome criteria
The patients’ records were analyzed in terms of the age at which the LM was diagnosed, birth history, sex, symptoms, symptom onset, type according to Olney’s classification, weight, and comorbid conditions. The feasibility of the STRIVE Hi anesthetic technique was evaluated according to the operation time, requirement for rescue oxygenation, and complications. Surgical outcomes, including resolution of symptoms, weight change, and complications, were also analyzed in patients who had undergone the operation. Weight percentile was analyzed using Wilcoxon’s signed-rank test with IBM SPSS ver. 22.0 (IBM Corp., Armonk, NY, USA).

RESULTS

Demographic and clinical data
A total of 29 LM patients were included in the study. The patients’ characteristics are summarized in Table 1. Twenty (68.9%) were female. Type I was the most common (n=13, 44.8%), fol-

Table 1. Clinical characteristics of the patients

| Variable                              | Number (%) |
|---------------------------------------|------------|
| Sex (female: male)                    | 20 (68.9):9 (31.1) |
| Type                                  |            |
| I                                     | 13 (44.8)  |
| II                                    | 10 (34.5)  |
| III                                   | 6 (20.7)   |
| Age of onset                          |            |
| At birth                              | 13 (44.8)  |
| >1 mo                                 | 4 (13.8)   |
| >2 mo                                 | 1 (0.3)    |
| Unknown                               | 11 (37.9)  |
| Symptom                               |            |
| Respiratory symptom                   |            |
| Stridor                               | 29 (100)   |
| Dyspnea with chest retraction         | 16 (55.2)  |
| Stridor worsening with upper respiratory infection or when crying, excited | 6 (20.7)   |
| Cyanosis, tachypnea                   | 4 (13.8)   |
| Feeding symptom                       |            |
| Small feeding amount, dysphagia due to respiratory distress | 12 (41.4)  |
| Tube feeding (nasogastric or gastrostomy tube) | 9 (31.0)   |
| Vomiting when feeding                 | 7 (24.1)   |
| Aspiration                            | 6 (20.7)   |
| Sleeping symptom                      |            |
| Snoring, apnea                        | 6 (20.7)   |

Table 2. Associated comorbidities

| Associated comorbidity                  | Number (%) |
|-----------------------------------------|------------|
| Congenital anomaly/syndrome/genetic disorder |            |
| Pierre Robin sequence                   | 1 (3.4)    |
| Down syndrome                           | 1 (3.4)    |
| Dandy-Walker (inversus) syndrome        | 1 (3.4)    |
| Trisomy 9                               | 1 (3.4)    |
| Cri du Chat variant                     | 1 (3.4)    |
| Mobius syndrome                         | 1 (3.4)    |
| Neurological disease                    |            |
| Seizure                                 | 4 (13.8)   |
| Developmental delay<sup>a</sup>         | 4 (13.8)   |
| Microcephaly, plagiocephaly<sup>b</sup> | 3 (10.3)   |
| Hypoxic ischemic encephalopathy        | 3 (10.3)   |
| Congenital hypotonia                    | 3 (10.3)   |
| Others<sup>c</sup>                      | 2 (6.9)    |
| Cardiac disease                         |            |
| ASD                                     | 9 (31.0)   |
| VSD                                     | 4 (13.8)   |
| PDA                                     | 2 (6.9)    |
| TOF                                     | 2 (6.9)    |
| TR                                      | 1 (3.4)    |
| PFO                                     | 1 (3.4)    |
| CMP                                     | 1 (3.4)    |
| Synchronous airway lesion               |            |
| Micrognathia, retrognathia              | 3 (10.3)   |
| Subglottic stenosis                     | 3 (10.3)   |
| Glossoptosis                            | 2 (6.9)    |
| Tracheomalacia, bronchomalacia          | 2 (6.9)    |
| Tongue base or vallecular cyst          | 2 (6.9)    |
| Vocal fold palsy                        | 2 (6.9)    |
| Cleft palate                            | 2 (6.9)    |
| Musculoskeletal                         |            |
| Developmental dysplasia of the hip      | 2 (6.9)    |
| Kyphoscoliosis, T-spine                 | 1 (3.4)    |
| A cyst in the filum terminale           | 1 (3.4)    |
| Partial agenesis of orbicularis oris muscle | 1 (3.4)  |
| Ophthalmologic                          |            |
| Retinal disorder                        | 1 (3.4)    |
| Unilateral esotropia with secondary ambyopia and bilateral optic neuropathy | 1 (3.4)   |
| Kidney                                  |            |
| Hydronephrosis                          | 3 (10.3)   |
| Others                                  |            |
| Mondini dysplasia, narrow BCNC (hearing loss) | 1 (3.4)   |
| Neonatal hyperbilirubinemia             | 4 (13.8)   |
| Disseminated intravascular coagulation, portal vein thrombosis | 1 (3.4)   |
| Neutropenia, aPTT prolongation          | 1 (3.4)    |
| BPD, RDS (hyaline membrane disease)     | 3 (10.3)   |

ASD, atrial septal defect; VSD, ventricular septal defect; PDA, patent ductus arteriosus; TOF, tetralogy of Fallot; TR, tricuspid regurgitation; PFO, patent foramen ovale; CMP, cardiomyopathy; BCNC, bony cochlear nerve canal; aPTT, activated partial thromboplastin time; BPD, bronchopulmonary dysplasia; RDS, respiratory distress syndrome.

<sup>a</sup>Language delay (n=1)+global developmental delay (n=3). <sup>b</sup>Plagiocephaly: microcephaly with simplified gyral pattern (n=3). <sup>c</sup>Cerebral infarct (n=1), brain cyst (n=1).
lowed by type II (n=10, 34.5%), and type III (n=6, 20.7%). About half of the patients (13, 44.8%) had symptoms at birth. All patients presented with stridor. Dyspnea with chest retrac-
tion was the second most common symptom, followed by re-
duced feeding amount and dysphagia due to respiratory distress. Of the 29 patients, 26 (89.7%) had comorbidities (Table 2). Among them, six patients (20.7%) had congenital syndromes 
associated with multiple anomalies. Two children (33.3%) with 
congenital anomalies/syndromes underwent supraglottoplasty. Of these, 11 (37.9%) had neurological disorders. The most com-
mon was seizure and developmental delay, and five (45.5%) 
underwent supraglottoplasty. Thirteen children (44.8%) had 
cardiac disease. The most common anomaly was atrial septal de-
fect. Eight (61.5%) of the patients with cardiac disease under-
went supraglottoplasty and two (15.4%) underwent heart sur-
gery. One (7.7%) underwent both procedures. SALs were found 
in 10 (34.5%) patients and four (40.0%) underwent supraglot-
toplasty. The most common SALs were micrognathia and retro-
gnathia. Fourteen patients (48.2%) had other comorbidities. Pre-
operatively, none had a patent tracheostoma.

Table 3. Perioperative characteristics of the patients who underwent surgical intervention

| No. | Sex | Age (mo) | Type | Comorbidity       | Simultaneous airway lesion | Operation                                              | Operation time (min) | Rescue oxygenation |
|-----|-----|----------|------|-------------------|-----------------------------|-------------------------------------------------------|---------------------|-------------------|
| 1   | M   | 3.1      | I    | G/N/S/O           | -                           | Microlaryngobronchoscopy+supraglottoplasty            | 35                  | -                 |
| 2   | F   | 9.3      | I    | N/S/O             | Glossoptosis                | Microlaryngobronchoscopy+supraglottoplasty           | 55                  | -                 |
| 3   | F   | 2.6      | I    | G/N/C/S           | uVFP, tracheomalacia, bronchomalacia | Microlaryngobronchoscopy   | 30                  | -                 |
| 4   | F   | 4.1      | I    | G/C/S/R           | Glossoptosis                | Microlaryngobronchoscopy+supraglottoplasty          | 145                 | Orotracheal intubation |
| 5   | F   | 3.4      | II   | G/N/S             | SGS, grade I                | Microlaryngobronchoscopy+supraglottoplasty          | 30                  | -                 |
| 6   | F   | 3.3      | II   | C/S/O             | SGS, grade II               | Microlaryngobronchoscopy+supraglottoplasty          | 66                  | -                 |
| 7   | F   | 3.9      | II   | O                 | -                           | Microlaryngobronchoscopy+supraglottoplasty          | 50                  | -                 |
| 8   | M   | 2.9      | II   | M/O               | -                           | Microlaryngobronchospy+supraglottoplasty            | 60                  | -                 |
| 9   | M   | 4.1      | II   | N/C/O             | -                           | Microlaryngobronchospy+supraglottoplasty            | 75                  | -                 |
| 10  | F   | 7.2      | II   | C/O               | -                           | Microlaryngobronchospy+supraglottoplasty            | 70                  | -                 |
| 11  | F   | 15.2     | II   | G/N/C/S           | Valacular cyst, SGS, grade II | Microlaryngobronchospy+supraglottoplasty+valacular cyst laser excision | 60                  | -                 |
| 12  | M   | 19.9     | II   | N/C/R             | -                           | Microlaryngobronchospy+supraglottoplasty            | 75                  | -                 |
| 13  | F   | 2.5      | III  | C                 | -                           | Microlaryngobronchospy+supraglottoplasty            | 140                 | -                 |
| 14  | F   | 19.0     | III  | N                 | -                           | Microlaryngobronchospy+supraglottoplasty            | 15                  | Orotracheal intubation (due to muscle relaxant application for severe trismus) |
| 15  | F   | 1.7      | III  | G/C/S             | Micrognathia, retrognathia, cleft palate, bronchomalacia | Microlaryngobronchospy | 60                  | Intermittent mask bagging |
| 16  | F   | 3.4      | III  | C/S/O             | Micrognathia, tongue base cyst | Microlaryngobronchospy+supraglottoplasty+tongue base cyst excision | 35                  | Orotracheal intubation |
| 17  | M   | 1.9      | III  | S                 | Micrognathia, glossoptosis, tongue base cyst | Microlaryngobronchospy+supraglottoplasty+tongue base cyst excision | 80                  | Orotracheal intubation |

N, neurological disease; S, synchronous airway lesion; O, ophthalmological; G, genetic disorders; C, cardiac disease; R, renal; O, others; M, musculoskeletal; uVFP, unilateral vocal fold palsy; SGS, subglottic stenosis.

*Operation time indicates the time interval for maintenance of anesthesia under the STRIVE Hi technique.
*As chest retraction was not alleviated after supraglottoplasty, the surgeon decided to perform tracheostomy after conversion to orotracheal intubation. There was no need for rescue oxygenation during supraglottoplasty. *Cases with planned tracheostomy.
Patients with surgical intervention
A total of 17 children (58.6%) underwent microlaryngobronchoscopy, 15 of whom continued to undergo supraglottoplasty under STRIVE Hi anesthesia (Table 3). Most type I patients did not require surgical intervention (4 of 13 patients required supraglottoplasty or tracheostomy, 30.8%). By contrast, most of the type II and III patients required surgery, including supraglottoplasty or tracheostomy (8/10, 80% vs. 5/6, 83.3%, respectively). Tracheostomy was planned for the remaining two patients without supraglottoplasty. Of these, one with micrognathia underwent tracheostomy (case 15), and the other patient with tracheomalacia and bronchomalacia did not undergo tracheostomy due to parental disagreement (case 3), and the symptoms improved slightly after 1 year of follow-up.

STRIVE Hi anesthesia maintenance and requirement for rescue oxygenation
The average age at the operation using the STRIVE Hi technique was 6 months (standard deviation [SD], 5.9 months; range, 1–20 months). Twelve patients (70.6%) completed the diagnostic procedure and surgical intervention without desaturation events (Fig. 2). The overall lowest SpO$_2$ was 75% (mean ± SD, 86.53% ± 12.17%; median, 88%) and the highest TcCO$_2$ was 75.2 mmHg (mean ± SD, 64.83 ± 10.70 mmHg; median, 69.2 mmHg). The mean self-ventilation maintenance time was 63.6 minutes (range, 15–145 minutes). The STRIVE Hi technique could not be maintained in one patient because of the requirement for muscle relaxation agents due to severe trismus. Desaturation events under the STRIVE Hi technique occurred in four cases, which ultimately led to conversion to conventional intubation in three cases, while the other was assisted with intermittent mask bagging (Table 3). The three patients in whom conversion to conventional intubation was performed had SALs, such as glossoptosis, tongue base cyst, and micrognathia. In one case, intermittent desaturation events occurred during the operation, which were assisted by intermittent mask bagging while maintaining the STRIVE Hi technique. In all cases, surgery was completed without any complications.

As the larynx could swell immediately after surgery, all patients were routinely administered epinephrine and inhaled salbutamol via a nebulizer. After supraglottoplasty, a high-flow nasal cannula was used in four patients (4/15, 28.6%) and eight patients (8/15, 57.1%) were monitored in the intensive care unit postoperatively. All patients were transferred to the general ward within 24 hours.

Surgical outcomes
Because the most of the patients were admitted to Department of Pediatrics, daily follow-up of symptoms was possible for 1 or 2 weeks after the operation. Symptoms usually improved on postoperative day 1 or 2 after the mucosal edema subsided. Symptom improvement was observed in 14 (93.3%) of the 15
The symptoms in LM patients are generally managed with conservative and medical therapy [4]. Despite the primarily spontaneously resolving nature of LM, 51.7% (15/29) of patients with LM and 61.5% (8/13) of patients with both LM and cardiac disease underwent supraglottoplasty at our center. This proportion is higher than in previous reports (12%–31% and 34%, respectively) [1,4,12-14].

The proportion of cases undergoing surgical treatment in our study was higher than that reported in previous studies because we had a higher proportion of patients with comorbidities and type II/III LM according to Onley’s classification. In previous studies, the rates of associated syndromes were 8%–20%, neurological disorders were 20%–45%, cardiac diseases were ~10%, and SAL were 7.5%–64% [1,15-17]. One of the most common features of SAL is the presence of other types of airway malacia, such as tracheomalacia and/or bronchomalacia (2%–48%) [1,3,11,15], and another common SAL is facial malformation (micrognathia and retrognathia, 10.3%). The presence of associated diseases increases the risk for negative outcomes after supraglottoplasty and the likelihood of performing tracheostomy [13,18]. In our study, the prevalence of comorbidities was similar to or higher than in previous studies (syndromes, neurological disorders, cardiac diseases, and SAL: 20%, 40%, 46.7%, and 36.7%, respectively). Another reason for the higher rate of surgical cases may have been the relatively high rates of type II and III LM in our cohort (type I, 43.3%; type II, 30%; type III, 23.3%). Olney et al. [1] reported that 57% of patients had type I LM, 15% had type II, 13% had type III, and 15 had combined types. According to the study, most patients with type I did not require surgery. By contrast, most type II/III LM patients required surgery, suggesting that type I may be a milder form than type II/III.

There are six possible anesthetic methods that can supply adequate oxygen during airway management: spontaneous breathing; positive pressure ventilation via an endotracheal tube; jet ventilation; apneic oxygenation using humidified high-flow nasal oxygen delivery systems or intermittent ventilation; cross-field ventilation; and extracorporeal membrane oxygenation (ECMO)/cardiopulmonary bypass (CPB) [19]. In the conventional endotracheal tube ventilation technique, the tube hides the subglottic airway and the surgical field. Sometimes, a tube can be intermittently withdrawn and excision can be performed while the child is apneic [2]. However, the process of inserting and removing the tube can cause airway trauma and prolong the operation time. The apneic technique with intermittent ventilation allows only approximately 3 minutes until further ventilation and oxygenation for a 3- to 4-year-old child [20,21]. In cases with associated conditions, such as cardiac, pulmonary, and airway lesions, which can affect cardiac output, the time to desaturation is likely to be even shorter [21,22]. The jet ventilation technique provides a tubeless surgical field. However, it has several disadvantages associated with the high airway pressure, such as an elevated risk for barotrauma (including pneumo-
rax, pneumomediastinum, and subcutaneous emphysema) and low reliability of end-tidal CO\(_2\) monitoring [19]. The cross-field ventilation technique requires tracheostomy, and ECMO/CPB is indicated in open surgery when manipulation of the heart and the great vessels is planned. The STRIVE Hi technique is a tubeless self-respiratory anesthesia method maintained intravenously, while additional oxygen and ventilation are supplied by a high-flow nasal cannula [9,23]. The high-flow nasal cannula allows oxygen delivery without the presence of any equipment in the surgical field and airway. In addition, it minimizes interruptions for rescue oxygenation even in settings of apnea or hyperventilation [24]. However, as mentioned, desaturation events occurred in four cases, which led to the requirement for oxygenation rescue [5]. Monitoring of ORI with SpO\(_2\) provides an early warning before oxygen saturation begins to decrease, which helps to avoid unanticipated hypoxia events [25]. By checking the ORI and SpO\(_2\) value, the anesthesiologists anticipated hypoxia, recognized the need to perform oxygenation rescue at the appropriate time, and finished preoxygenation without hypoxia. All oxygen rescue processes were continued with sufficient multidisciplinary communication. When an anesthesiologist notified the team of signs of desaturation, the operation was stopped and temporary tracheal intubation was performed through a Benjamin-Lindholm laryngoscope. Likewise, continuous CO\(_2\) monitoring is essential because the elevation of CO\(_2\) needs correction during the operation. Mask bagging was performed by an anesthesiologist. As a result, all operations were completed without severe adverse events while intubation was performed in three cases and intermittent mask bagging was performed in one case (Table 3). In another case in which the patient did not have SAL, oxygen saturation was stable with the STRIVE Hi technique. However, a muscle relaxant had to be applied because the surgical site could not be exposed due to severe trismus, which led to the loss of self-respiration and conversion to conventional intubation. After experiencing several cases with the STRIVE Hi technique, we would suggest that this technique is contraindicated for patients with a skull base fracture or choanal atresia. In addition, poor exposure of the surgical field, such as in cases of micrognathia, is a relative contraindication for transoral supraglottoplasty under the STRIVE Hi technique.

During the operation using the STRIVE Hi technique, temperature and humidity were maintained uniformly for all patients at levels that were high enough to prevent feelings of dryness in the nasal cavity of patients and to promote tolerance for the high-flow rate [26]. In addition, higher temperature and humidity than the gas supplied in conventional anesthesia can decrease intraoperative oxygen demand by maintaining body temperature higher [27], and postoperative lung complications may decrease by maintaining the mucociliary clearance of the airway [28]. The flow rate of O\(_2\) was 2 L/min/kg according to previous reports and experiences at our institution [5]. There were no complications associated with the use of the STRIVE Hi technique such as pneumothorax, gastrointestinal air, or difficulty obtaining a seal during masking. There was no fire when we used a CO\(_2\) laser under the STRIVE Hi technique, probably due to the lack of a medium to support ignition such as an endotracheal tube.

LM is suspected based on a typical clinical history, but the diagnosis is confirmed by a flexible fiberoptic exam in conscious patients [10]. Sivan et al. [29] reported that anesthesia/sedation-aided diagnosis may sometimes be more accurate than conscious examination due to rapid laryngeal motion and lack of patient cooperation. Patients with severe LM may be at greater risk for SALs causing multilevel airway obstruction [2,16]. This supports the importance of conducting microlaryngobronchoscopy for a multilevel airway assessment before supraglottoplasty. Using the STRIVE Hi technique, we could secure the surgeon’s view without blocking the endotracheal tube and perform accurate preoperative evaluation of airway dynamics. During supraglottoplasty, movements of the supraglottis did not hinder the surgical procedure as the surgical site was immobilized with cup forceps or alligator forceps. The postoperative outcome could be checked in real-time during supraglottoplasty, and the surgeon could decide whether the patient required tracheostomy before leaving the operating room.

The success rate of supraglottoplasty is approximately 94%, and it has a low complication rate. In this study, using the STRIVE Hi technique, 93.3% (14/15) of patients who underwent supraglottoplasty showed symptom improvement without severe complications. Children with LM are at significant risk for insufficient weight gain due to an increased workload while breathing and increased energy expenditure [3]. The most frequent reasons for surgical intervention are worsening airway symptoms and failure to thrive [30]. Similarly, the most important indicator of the improvement of LM in infants is appropriate weight gain [3]. Our patients showed significant changes in weight percentile after supraglottoplasty (\(P=0.026\)).

Patients with LM referred to our tertiary hospital had higher comorbidity rates than previously reported. Supraglottoplasty is a safe and effective treatment for most cases of severe LM. The STRIVE Hi technique ensures that the surgeon has an appropriate view for accurate diagnosis and ease of operation, and enables evaluation of the immediate postoperative respiratory status and results. Type III LM patients with micrognathia or glossoptosis may have a higher likelihood of requiring rescue oxygenation during the STRIVE Hi technique, while the presence of multiple comorbidities seemed to have no effect on the surgical success of supraglottoplasty.

**CONFLICT OF INTEREST**

No potential conflict of interest relevant to this article was reported.
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ORCID

Hyunkyung Cha https://orcid.org/0000-0002-3462-3789
Doh Young Lee https://orcid.org/0000-0003-1590-8559
Eun-Hee Kim https://orcid.org/0000-0002-8384-8191
Young-Eun Jang https://orcid.org/0000-0002-7511-4104
Hee-Soo Kim https://orcid.org/0000-0002-2661-7944
Seong Keun Kwon https://orcid.org/0000-0001-9218-7666

AUTHOR CONTRIBUTIONS

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