A comparison of supraglottic devices in pediatric patients

Senthil G Krishna1,2
Faizaan Syed1
Mohammed Hakim1
Mumin Hakim3
Dmitry Tumin1
Giorgio C Veneziano1,2
Joseph D Tobias1,2

1Department of Anesthesiology & Pain Medicine, Nationwide Children’s Hospital, Columbus, OH, USA; 2Department of Anesthesiology & Pain Medicine, The Ohio State University College of Medicine, Columbus, OH, USA; 3Department of Surgery, Montefiore Medical Centre, Bronx, NY, USA

Background: When managing patients with a difficult airway, supraglottic airways (SGAs) have been used as rescue devices or to serve as a conduit for endotracheal intubation. The current study compares various clinical outcomes, including the bronchoscopic view of the glottis when using 2 SGAs, the Air-Q® laryngeal mask airway (LMA) and the i-gel® SGA, in pediatric patients.

Methods: Patients ≤18 years of age were prospectively randomized to receive either the Air-Q® LMA or the i-gel® SGA. Following SGA placement, a flexible fiberoptic bronchoscope was inserted through the SGA to visualize the glottis. Time taken to obtain the bronchoscopic view and place the SGA, and the ability to seal the airway at 20 cmH2O were compared. The bronchoscopic view obtained was graded as follows: 1) glottic aperture seen completely; 2) glottic aperture seen partially with visual obstruction <50%; 3) glottic aperture seen, but visual obstruction >50%; and 4) glottic aperture not seen.

Results: Fifty patients were enrolled and 48 (22/26 male/female) were included in the analysis. Median age was 13 years (IQR: 7, 16) and median weight was 49 kg (IQR: 25, 70). The Air-Q® LMA and i-gel® SGA groups did not differ. The time required for successful placement of the SGA, the time required for bronchoscopic view, and the quality of bronchoscopic view through the Air-Q® LMA and the i-gel® SGA did not differ.

Discussion: The time required for successful placement of the SGA, the time required for bronchoscopic view, and the quality of bronchoscopic view through the Air-Q® LMA and the i-gel® SGA did not differ.

Keywords: supraglottic airways, bronchoscopic view, glottic aperture

Introduction

A difficult airway is generally defined as a challenging situation where there is an inability or difficulty in establishing bag-mask ventilation or difficulty with conventional direct laryngoscopy.1 This presents an even greater challenge in children compared with adults due to certain anatomical variations and physiologic considerations. Hence, in children, there may be less time available resulting in an increased incidence of clinical deterioration prior to re-establishing oxygenation and ventilation.2 The most common complication associated with management of the difficult airway is transient hypoxemia; however, the most common severe complication in a failed airway is cardiac arrest.3 Supraglottic airways (SGAs) have been used successfully in the management of difficult airways in either re-establishing and maintaining oxygenation and ventilation or as a conduit for fiberoptic-guided endotracheal intubation.1 SGAs can also
overcome upper airway obstruction and maintain an airway without the need for endotracheal intubation.

Although various SGAs have been used as rescue airway devices, data on the relative performance of SGAs remain limited in terms of which SGA may be a preferred choice in pediatric patients with a difficult airway. Jagannathan et al compared the Air-Q® laryngeal mask airway (LMA) with the i-gel® SGA and found that they both were equally effective in providing an adequate fiberoptic view when placed by trainees, but more problems were encountered during removal of the i-gel® SGA, including dislodgement of the endotracheal tube (ETT). By contrast, Kim et al reported that the Air-Q® LMA provided an improved fiberoptic view over the i-gel® SGA although the i-gel® SGA was easier to place.

Given this ambiguity over the relative utility of the 2 devices in the aforementioned studies, and variability in the results using other SGAs, we chose to compare the ease of placement, adequacy of ventilation, and the bronchoscopic fiberoptic view through the internal lumen of the Air-Q® LMA and the i-gel® SGA in pediatric patients. By examining the fiberoptic view through these 2 devices, we sought to determine which SGA would provide a clear pathway to the glottic aperture and therefore, potentially provide an easier path for endotracheal intubation.

Methods

The study was approved by the Institutional Review Board (IRB) of Nationwide Children’s Hospital with a waiver of individual consent (IRB15-00486) as both types of SGAs were accepted standard clinical practice. The trial was registered with ClinicalTrials.gov (NCT02532465). American Society of Anesthesiologists grade 1 or 2 patients who were ≤18 years of age and scheduled to receive an SGA as part of standard anesthetic care for elective surgery were enrolled and randomized to receive 1 of 2 types of SGA devices: the Air-Q® LMA or the i-gel® SGA. Patients with a known or suspected difficult airway or a history of prior difficult placement of an SGA were excluded from the study. The SGA chosen by randomization was revealed to the investigator and the anesthetic team immediately prior to device placement. The size of the SGA was determined according to the manufacturer’s recommendations based on the patient’s weight.

The induction of anesthesia was achieved with 70% N₂O in O₂ and sevoflurane. Following placement of the intravenous cannula, N₂O administration was terminated and anesthesia was supplemented with intravenous propofol (2 mg/kg) and fentanyl (2 µg/kg). The SGA was placed using a standard midline technique with neck flexion, head extension, mouth opening, and anterior displacement of the tongue. The time taken to place the SGA, defined as time from the removal of the anesthesia mask to reappearance of capnographic activity, was noted. Ease of insertion was assessed by the provider placing the SGA using a previously described subjective grading score of 1–4 for SGAs (1= no resistance; 2= mild resistance; 3= moderate resistance; and 4= inability to insert the device). If the study SGA placement was unsuccessful, the airway was secured using an Ambu® LMA. Following satisfactory placement of the SGA, air was added to the cuff of the SGA if needed to achieve adequate chest movement with positive pressure ventilation. The ability to seal the airway at 20 cmH₂O without an audible leak was noted using the manometric stability test with a fresh gas flow at 3 L/min. Optimal position of the SGA was confirmed by auscultation and generation of an adequate tidal volume. If the ventilation was judged to be suboptimal by the clinical team, the airway was rescued using an Ambu® LMA.

After confirmation of adequate placement and ventilation, a flexible fiberoptic bronchoscope was inserted through the stem of the supraglottic device to visualize the glottic aperture. The time taken for bronchoscopy, defined as time from disconnection of the anesthetic circuit from the SGA to first visualization of the glottic aperture, was recorded. The bronchoscopic view obtained via the intubating SGA was graded using our scale developed from previously used scales: Grade 1= glottic aperture seen completely without any obstruction; Grade 2= glottic aperture seen only partially but visual obstruction is <50%; Grade 3= glottic aperture barely seen and visual obstruction is >50%; and Grade 4= glottic aperture not seen (Table 1). Complications, including bleeding, oxygen desaturation, laryngospasm, and bronchospasm, were also noted.

Categorical data were presented as counts with percentages and compared using Fisher’s exact tests or chi-squared

### Table 1 Grading of fiberoptic view of glottic aperture through supraglottic device

<table>
<thead>
<tr>
<th>Glottic view grade</th>
<th>Description</th>
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<tr>
<td>1</td>
<td>Glottic aperture seen completely without any obstruction</td>
</tr>
<tr>
<td>2</td>
<td>Glottic aperture seen only partially but visual obstruction is &lt;50%</td>
</tr>
<tr>
<td>3</td>
<td>Glottic aperture barely seen and visual obstruction is &gt;50%</td>
</tr>
<tr>
<td>4</td>
<td>Glottic aperture invisible</td>
</tr>
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tests, as appropriate. The primary analysis compared the fiberoptic view through the 2 different supraglottic devices. The percentage of acceptable or good views (grade 1 or 2) was compared with poor view (grade 3 or 4). Continuous data were presented as median with IQR and compared using rank-sum tests. Study enrollment was planned for 50 patients according to availability of study staff, and no a priori power analysis was performed. Data analysis was performed in Stata/IC 14.2 (StataCorp, LP, College Station, TX, USA) and 2-tailed \( P<0.05 \) was considered statistically significant.

**Results**

Fifty patients were enrolled in the study, with 25 randomized to each group. Ages ranged from 1 to 18 years (median: 13 years, IQR: 7, 16) and weight ranged from 11 to 106 kg (median: 49 kg, IQR: 25, 70). The size of the Air-Q® LMA varied from 1.5 to 4.5 and the size of the i-gel® SGA ranged from 2 to 5. Demographic characteristics for the 2 groups are summarized in Table 2. One patient in the Air-Q® LMA group was excluded due to missing study data. Except for 1 patient in the Air-Q® LMA group where 2 attempts were required, all SGAs were inserted on the first attempt. One patient in the i-gel® SGA group required removal of the SGA device and placement of an Ambu® LMA due to inadequate ventilation. In all other patients, placement of the SGA was judged as easy (1=no resistance). In one other case, the i-gel® SGA was replaced with an Ambu® LMA after completion of data collection due to loss of ventilation. In another patient, the Air-Q® LMA was replaced with an Ambu® LMA after study data were obtained due to problems with placement of the esophagogastroduodenoscopy scope.

An acceptable fiberoptic view (grade 1 or 2) was attained in 20 of 24 cases in the Air-Q® LMA group and 21 of 23 in the i-gel® SGA one (\( P=0.666; 1 \) case in the i-gel® SGA group had missing data on this outcome). Time taken to place the SGA did not significantly vary between the Air-Q® LMA and i-gel® SGA (median of 19 [IQR: 16, 20] vs 21 [IQR: 16, 29] seconds; 95% CI of difference in medians: −2 to 7; \( P=0.331 \)). Likewise, time to achieve fiberoptic view of the glottis was similar between the 2 groups (median of 25 [IQR: 20, 30] vs 21 [IQR: 17, 40] seconds; 95% CI of difference in medians: −9 to 8; \( P=0.489 \)). Eight Air-Q® LMAs and 6 i-gel® SGAs sealed the airway at 20 cmH\(_2\)O (Table 3).

**Discussion**

Adequate airway control with effective oxygenation and ventilation remain integral aspects of airway intervention regardless of the device used. Compared with adults, children have increased oxygen consumption coupled with a lower oxygen reserve thereby limiting their tolerance to apnea, and increasing the incidence of hypoxemia and its cardiovascular and hemodynamic sequelae.\(^17\) SGAs have proven to be extremely useful in these clinical scenarios, both as a means to restore oxygenation and ventilation as well as serving a conduit for the fiberoptic-guided endotracheal intubation.\(^5,6\)

The current study compared the performance of 2 commonly used SGAs; the Air-Q® LMA and the i-gel® SGA.

Modifications of the Air-Q® LMA to facilitate its use for device-guided endotracheal intubation include a shorter and curved shaft, no epiglottic grill, and a removable 15 mm airway adapter.\(^18,19\) However, we hypothesized that the design of the i-gel® SGA, including elimination of the need to inflate the cuff, integrated bite block, and the wide oval stem may improve in vivo stabilization, prevent axial rotation, and facilitate obtaining a bronchoscopic view of the glottis when compared with the Air-Q® LMA. Similar to the previous studies,\(^5–13\) as our goal was to objectively evaluate SGA use for fiberoptic-guided endotracheal intubation, we chose to use the fiberoptic view of the glottic aperture as one of the primary outcome measure to compare these 2 SGAs. Given concerns with the complexity of the grading scale\(^15,16\) and the design differences of the 2 devices, the current study used a simplified version to allow head-to-head comparison of the 2 SGAs based on the clinically relevant glottic view obtained with the device.\(^5,6\)

Previous studies directly comparing these 2 SGAs have provided contradictory results.\(^5,6\) Jagannathan et al compared the performance of the i-gel® SGA with the Air-Q® LMA in a cohort of 96 children, ranging in age from 1 month to 6 years with anatomically normal airways.\(^3\) The authors noted that the 2 devices were equally effective as conduits for fiberoptic-guided endotracheal intubation even when performed by trainees with limited prior experience. They determined that use of the i-gel® SGA resulted in a faster

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**Table 2 Characteristics of patients enrolled in each study group**

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<tr>
<th>Characteristic</th>
<th>Air-Q® LMA</th>
<th>i-gel® SGA</th>
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<tr>
<td>Male</td>
<td>12 (50%)</td>
<td>10 (42%)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>14 (6, 16)</td>
<td>12 (7, 16)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>50 (22, 70)</td>
<td>49 (26, 69)</td>
</tr>
<tr>
<td>SGA size</td>
<td>3.5 (2, 3.5)</td>
<td>3 (2.5, 4)</td>
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time to first glottic view, time to carinal view, and time to successful endotracheal intubation. However, the i-gel® SGA was associated with more problems during removal with the ETT in place, including dislodgement, need to hold downward traction, pilot balloon breakage, or inadvertent extubation. However, Kim et al in a cohort of 80 children reported that the i-gel® SGA was easier to insert with higher oropharyngeal leak pressures and lower frequencies of gastric insufflation compared with the Air-Q® LMA. In contrast to what was noted by Jagannathan et al, they reported that the Air-Q® LMA provided a better fiberoptic view of the glottis than the i-gel® SGA.6

We found no clinically significant differences in the performance of the 2 devices for the tested parameters, which included their placement, function as an SGA during intraoperative care, or the fiberoptic view of the glottis that was achieved. Both SGAs were easy to place and provided a similar and acceptable view of the laryngeal structures with no difference in time required to achieve placement or glottis view. Furthermore, both devices performed similarly with respect to sealing the airway at 20 cmH2O. Minor airway events, which required replacement of the devices with the standard LMA that we use in clinical practice were noted in 2 patients with the i-gel® SGA (once immediately following placement and once during subsequent use) and in 1 patient with the Air-Q® LMA. Anecdotally, anesthesia providers in our department perceived the Air-Q® LMA to be easier to use than the i-gel® SGA, but this subjective preference was not quantified in the present study, and not reflected in the objective measures of performance. Furthermore, our clinical practice prior to the study included sole use of the Ambu® or Air-Q® LMA thereby resulting in significant clinical experience with these devices as opposed to little or no prior experience with the i-gel® SGA.

With comparable performance between the devices based on our findings, the decision to favor one over the other may be influenced by other clinical factors, including the ease of advancement of the fiberoptic bronchoscope, placement of the ETT, and removal of the intubating SGA. While Jagannathan et al found no difference between the SGAs in terms of navigation of the fiberoptic bronchoscope or advancing the ETT through the device, they reported clinical problems with removal of the i-gel® SGA, including dislodgement of the ETT (see aforementioned). The particular feature of ease of removal of the SGA after placement of the ETT may be one factor that differentiates these 2 devices. The Air-Q® LMA, which is designed with a large inner diameter, relatively shorter stem and detachable 15 mm connector piece may facilitate removal of the device following endotracheal intubation. However, this particular outcome was not specifically studied in our patient population.

Limitations of our current study include that it was not limited to a younger patient population as in previous studies, but rather enrolled the entire spectrum of pediatric ages. Although we acknowledge that younger children with associated difficult airways are relatively more challenging, we designed our study to be a representation of the patient population that we encounter in our routine clinical practice.

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the ease of placement of an ETT through the SGA and the ease of subsequent removal of SGA. Finally, our study was conducted with a convenience sample and therefore may have had limited statistical power to compare rare events between the 2 SGA types, including subsequent intraoperative performance in various clinical scenarios and varying surgical procedures. With these caveats in mind, the current study adds further data comparing 2 of the most commonly used SGAs in the pediatric-aged patient. We noted that there were no clinically significant differences in the performance of the 2 devices in the tested parameters. Both devices performed equally well despite the fact that the clinical staff had limited previous experience with the i-gel® SGA, which further demonstrates that clinical expertise with placement of SGAs can be rapidly acquired. Additional studies are needed to evaluate whether previously noted problems identified during removal of the i-gel® SGA after endotracheal intubation make the Air-Q® LMA the preferred choice of SGA for the management of the difficult airway in children.

Disclosure
The authors report no conflicts of interest in this work.

References