Use of ultrasonography in confirming the malposition of the i-gel in pediatric patients: a prospective observational study

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Use of ultrasonography in confirming the malposition of the i-gel in pediatric patients: a prospective observational study

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Background/aim: This study was designed to observe and compare the performance of fiberoptic laryngeal (FOL) view, leakage test (LT) and ultrasonography (USG) usage in detecting i-gel position in pediatric patients.

Materials and methods: One hundred ten consecutive children were included in this single-armed prospective observational study. After anesthetic induction, i-gel placement confirmed using FOL and LT was evaluated using USG in three planes. According to our scoring system, acceptable and unacceptable grades (FOL, LT and USG) were determined to describe placement. Sensitivity and specificity were determined by comparing USG performance with the other two tests.

Results: Ultrasonography was found a sensitivity of 20% positive predictive value (PPV) for unacceptable i-gel placement according to FOL grade and a sensitivity of 37.04% with a 100% PPV according to LT grade. USG was found a specificity of 91.84% negative predictive value (NPV) of 91.84% for acceptable i-gel placement according to FOL grade and the NPV specificity of 100% with a 82.65% NPV according to LT grade.

Conclusion: Ultrasonography demonstrated a very good diagnostic performance in the detection of optimal i-gel placement according to both FOL and LT. However, both FOL and LT showed poor diagnostic performance compared to USG in demonstrating i-gel malposition.

Key words: Airway management, fiberoptic examination, leakage test, ultrasonography

1. Introduction
The use of i-gel, as a relatively new supraglottic airway device (SAD) released in 2007, has become popular for children undergoing surgery with general anesthesia because it does not require muscle relaxation. It is important to place an i-gel in the optimal position to ensure adequate ventilation and avoid complications such as mucosal injury, glottic ptosis, and possible aspiration and gastric insufflation [1,2]. Successful insertion is usually assessed using a capnogram with an end-tidal carbon dioxide (EtCO₂) value, appropriate chest elevation, no more than 20 cmH₂O peak inspiratory pressure and the absence of audible oropharyngeal leakage [2]. Children have broad tongues and a drooping epiglottis. Also, their larynx is higher and more anterior than in adults. This difference in airway anatomy may affect the correct placement of SADs [3]. Although fiberoptic laryngeal (FOL) view is considered the preferred verification tool for direct visualization of the glottis, it has also been reported to be a superior technique for detecting the malposition of SADs [4–6]. Some studies of SAD position using FOB have reported that a proportion of children need repositioning of smaller SADs (12.8%–49%) despite adequate ventilation [2,7,8]. Ultrasonography (USG), which is a noninvasive, simple and portable technology, has recently come into use in evaluating airway management even in upper airway anatomy impaired due to pathology or trauma [9,10]. However, although several publications on ultrasonography confirming the optimal location of various SADs have been reported recently, USG publications examining its malposition have not been reported [11,12].

The aim of this study was to compare the FOL image and leakage test (LT) values used in the placement of i-gels with USG findings in pediatric patients and to demonstrate the performance of USG in confirming i-gel malposition.
2. Methods
This single-arm prospective observational study was reviewed and approved by the Selçuk University Faculty of Medicine the Local Ethics Committee (14.10.2020.2020/19) and recorded in the Clinical Trials Registry (ClinicalTrials.gov Identifier: NCT04652466). After obtaining written informed consent from the parents of the participants, 110 patients with American Society of Anesthesiologists (ASA) status I-II undergoing elective pediatric surgery and had i-gels placed were enrolled in the study. Patients with head and neck anatomic malformations and airway congenital defects were excluded from the study.

Midazolam 0.5 mg kg\(^{-1}\) was administered orally 20 min before surgery for routine premedication to all patients. The patients were placed on the operating table in the supine position. All patients were monitored with standard monitoring (electrocardiogram, noninvasive blood pressure, oxygen saturation, \(\text{ETCO}_2\), peak airway pressure), after a minimum of 2 min of preoxygenation, inhalational induction was performed with 6%–7% sevoflurane at the beginning of general anesthesia induction. IV access was provided, and according to standard procedures, propofol 3.5 mg/kg, fentanyl 1 mcg/kg iv and remifentanil 0.1 mcg/kg/min infusion was administered. The i-gel size was selected according to the manufacturer's recommendations. Adequate anesthesia depth was confirmed by loss of eyelash reflex, symmetrical small pupils, and absence of swallowing, and i-gel was gently guided along the hard palate by the first anesthesiologist until resistance was felt, with the i-gel (Intersurgical Ltd., Wokingham, Berkshire, UK) opening facing the applicator. Patients were mechanically ventilated with an \(\text{ETCO}_2\) value between 35 and 45 mmHg with volume-controlled ventilation, at 3 L/min of fresh gas flow, 6–8 ml/L tidal volume, and 18 to 24 breaths/min frequency. After 5 ventilator cycles, \(P_{\text{peak}}\) was recorded. If the patient did not tolerate the insertion of the i-gel, an extra dose of fentanyl up to a maximum of 0.5 mcg/kg was administered. However, if the insertion attempt was still not tolerated, or if the capnography curve was not obtained and there was an audible leakage, the technique was considered to be unsuccessful, and another supraglottic SAD was then placed in the airway. If this change was not tolerated, endotracheal intubation was initiated by administering a neuromuscular blocker and these were excluded. All patients were evaluated by a total of four experienced independent anesthetists who performed intubation, FOL, LT, and USG examinations.

2.1. Fiberoptic laryngoscopy examination
A fiberoptic laryngoscope was passed between the Y-connector and the i-gel to obtain the FOL by the second anesthesiologist. The image acquired by the FOL advanced from i-gel was categorized as described by Campbell et al. [13]. The optimal placement was defined as epiglottis visibility of less than 50% (Figures 1A–1C) where the epiglottis did not enter the airway through the i-gel; acceptable placement was recorded as FOL-A (fiberoptic laryngoscopy acceptable). Categories E and F were defined as more than 50% epiglottis visibility where the epiglottis entered the airway; unacceptable insertion was recorded as FOL-U (fiberoptic laryngoscopy unacceptable) (Figures 1D and 1E) (Table 1).

2.2. Leakage test (LT) examination
LT was determined by temporarily ceasing ventilation and closing the adjustable pressure-limiting valve with a fresh gas flow of 3 L min\(^{-1}\) until the airway pressure reached a steady state and a leakage sound was heard by the third anesthesiologist. The leakage pressure was not allowed to exceed 40 cm \(H_2O\). The LT grading system was based on the study of Theiler et al. [14]. Acceptable LT (LT-A) was recorded as peak inspiratory pressure (PIP) ≤ 26 cm H2O, bilateral chest lift and square wave capnogram formation, no square wave capnogram, an audible leakage. PIP ≥ 27 cm H\(_2\)O were recorded as unacceptable LT (LT-U) (Table 1).

2.3. Evaluation of i-gel placement using USG
USG examination of the neck was performed by the fourth anesthesiologist using an Esaote 8-18–MHz linear

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**Figure 1.** Fiberoptic appearance of the glottis: Percentage and categories of glottic opening covered by the epiglottis. A–C indicates that the epiglottis visibility is less than 50% and the epiglottis is positioned correctly. D and E indicates that the epiglottis visibility is more than 50% and the epiglottis is misplaced.
Table 1. Grading according to FOL, USG and LT.

<table>
<thead>
<tr>
<th>FOL grade</th>
<th>USG grade</th>
<th>Leakage test grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>FOL-A</td>
<td>USG-A</td>
<td>LT-A</td>
</tr>
<tr>
<td>FOL-U</td>
<td>USG-U</td>
<td>LT-U</td>
</tr>
<tr>
<td>The categories A-B-C</td>
<td>The scores 0–1</td>
<td>Leakage test ≤ 26 cmH₂O</td>
</tr>
<tr>
<td>The categories D-E</td>
<td>The scores 2–5</td>
<td>Leakage test ≥ 27 cmH₂O</td>
</tr>
</tbody>
</table>


2.4. Statistical analysis

The study by Kim et al. [11] on the difference between USG and FOL in the incidence of i-gel malposition was accepted as a reference. The incidence of i-gel malposition with FOL was 50% and 43% on USG. Assuming a 5% significance level (α = 0.05) and power of 80% (= 0.20) to detect a 7% absolute difference in the incidence of i-gel malposition with USG and FOL, a sample size of 102 patients was required. The sample size allowing for possible data loss was allocated as 110 patients.

All statistical analyses were performed using the online R 3.6.0 program (https://www.r-project.org). Data are presented as mean ± standard deviation (range: minimum-maximum) or numbers (n) and percentages (%). Kappa (κ) coefficient analysis was used to test the agreement between FOL, USG, and LT grades. The diagnostic performance was tested with sensitivity, specificity, positive and negative predictive value (PPV and NPV, respectively), and accuracy. The sensitivity was the percentage of USG grades 2 to 5 and LT grades higher than ≥ 27 that were correctly identified as i-gel malposition grades 4 to 5 using FOL. The specificity was the percentage of USG grades 0 to 1, and LT grades lower than ≤ 26 that were correctly identified as i-gel malposition grades 1 to 3. The PPV was the percentage of i-gel placement grades 4 to 5 when USG arytenoid image grades were 3 to 5, and LT grades were higher than ≥ 27. The NPV was the percentage of i-gel placements 1 to 3 when the USG arytenoid image was 0 to 1 and LT grades were lower than ≤ 26. Accuracy was the percentage of concordance between 2 devices in terms of grades. The confidence interval of the abovementioned parameters was calculated using the Clopper–Pearson method. Chi-square and Z-tests were used to compare the rate ratios of malposition between FOL, USG, and LT methods. Moreover, the McNemar test was used to compare the proportions of the methods. A p-value of less than 0.05 was considered statistically significant.

3. Results

A total of 110 patients underwent elective pediatric surgery between February 2021 and December 2020. A total of 2 patients were excluded for not meeting inclusion criteria. Thus, 108 patients were enrolled. Detailed information on enrollment of patients into the study is depicted in the CONSORT flow diagram in Figure 3. The demographic and clinical characteristics of the children are given in Table 2. A total of 108 children who had a mean age of 3.64 ± 2.29 (range, 1–10) years, mean BMI of 15.23 ± 3.05 (range, 3.79–25.14), were included in the study. The i-gel was successfully inserted at the first attempt in 98 (90.7%), at the second attempt in nine (8.3%), and at the third attempt in one (0.9%) child. The mean EtCO₂ was 39.82 ± 4.43 (range, 30–45), the mean SpO₂ was 98.77 ± 1.09.
There was no significant difference in Ppeak between the FOL grades ($p = 0.144$), but the mean Ppeak was higher in unacceptably positioned i-gels compared with acceptably positioned i-gels in both USG ($16.20 \pm 4.87 \text{ vs. } 11.60 \pm 2.66$, $p < 0.001$) and LT ($14.52 \pm 4.11 \text{ vs. } 11.19 \pm 2.32$, $p < 0.001$) grades (Table 2).

In the LT, 81 out of 108 (75%, 95% CI: 65.75%–82.83%) children whose LT scores were lower than 28 were found to be acceptable, and 27 (25%, 95% CI: 17.17%–34.25%) children whose LT score was $\geq 27$ were found to be unacceptable. The mean LT score of the children was $24.51 \pm 6.40$ (range, 13–39) (Table 3).

In the USG examination grade, 86 children (79.6%) were found to be grade 0, 12 (11.1%) grade 1, 7 (6.5%) grade 2, and 3 (2.8%) grade 3. USG grade $> 1$ was seen in 10 out of 108 children (9.26%, 95% CI: 4.53%–16.37%) in whom i-gel placement was defined as unacceptable, and USG grade $\leq 1$ was seen 98 (90.74%, 95% CI: 83.63%–95.74%) children, where i-gel was defined as acceptable (Table 3).

The rate of i-gel displacement was similar between FOL grades and USG grades ($p > 0.999$), but this incidence was lower in USG grade compared with LT grade ($p = 0.022$) (Table 3).

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The rate of i-gel displacement was similar between FOL grades and USG grades ($p > 0.999$), but this incidence was lower in USG grade compared with LT grade ($p = 0.022$) (Table 3).

Ninety out of 108 children had acceptable USG and FOL grades in i-gel placement. Two out of 108 children had acceptable USG and FOL grades in i-gel placement. Eighty-one out of 108 children had acceptable USG and LT grades in i-gel placement. Ten out of 108 children had acceptable USG and FOL grades in i-gel placement. Of the 98 i-gel placements that were positioned acceptably according to the FOL grade, 90 were successfully positioned using USG. In addition, of the 10 unacceptable
i-gel placements that were positioned according to FOL grade, two were successfully positioned using USG. Of the 81 acceptable i-gel placements that were positioned according to FOL grade, 81 were successfully positioned using USG. Of the 27 unacceptable i-gel placements that were positioned according to the FOL grade, 10 were successfully positioned using USG (Table 4).

Regarding the diagnostic performance of USG in terms of FOL and LT grade, USG was found to have a sensitivity of 20% (2.52%–55.61%), with a PPV of 20% (5.77%–50.50%) to detect an unacceptable i-gel placement according to FOL grade. The specificity was 91.84% (84.5%–96.41%), with an NPV of 91.84% (89.14%–93.91%) to detect an acceptable i-gel placement according to FOL grade. The accuracy was 85.19% (77.06%–91.29%). USG was found to have a sensitivity of 37.04% (19.40%–57.63%), with a PPV of 100% (69.2%–100%) to detect an unacceptable i-gel placement according to LT grade. The specificity was 100% (95.55%–100%), with an NPV of 82.65% (78.11%–86.42%) to detect an acceptable i-gel placement according to LT grade. The accuracy was 84.26% (76%–90.55%). The kappa value was 0.118 ± 0.096 between USG and FOL grade, and there was no statistically significant agreement between USG and FOL grade ($Z = 1.23, p = 0.218$). However, the kappa value was 0.468 ± 0.081, which suggests a moderate agreement between USG and LT grade, and this kappa value is significantly different from zero ($Z = 5.75, p < 0.001$) (Table 4).

4. Discussion
Fiberoptic laryngoscope view and LT grades were compared with USG grades in i-gel positioning. It was shown that USG was compatible with FOL and LT in showing the optimal i-gel placement, but its diagnostic performance was poor for malpositioned i-gels.

Although the most commonly used test to evaluate the position of SADs is the LT, its use in cases of laryngospasm or bronchospasm may be restricted, regardless of the position of the SADs in the oropharynx [14]. Therefore, imaging the airway with fiberoptic is the gold standard in confirming the location of SADs and avoiding airway-related adverse events [16].

However, although FOL assessment is an important assessment tool, it may not be used routinely because it
is technically demanding and an invasive procedure. Recently, there has been a great interest in the use of USG in airway management, which allows uninterrupted airway management as well as being safe, fast, portable, and repeatable. Previously limited to airway evaluation, it now finds use in real-time dynamic airway management [9,10].

The first report on USG evaluation for proper placement of SADs was published by Kim et al. In their study [11], they compared USG images of positional changes in arytenoids before and after insertion of classical LMA in 100 pediatric patients. USG was found to have a sensitivity of 93% and specificity of 82% (accuracy was 87% to detect a rotated LMA) [16]. In the LMA study of Zhou et al. clinical findings, FOL and USG were compared, and it was suggested that USG evaluation was superior to other techniques to confirm proper LMA placement [17]. In the last few years, several studies have been conducted in which USG verification of the position of SADs was evaluated and compared with other methods, and USG was found as an appropriate tool for confirming SAD placement [15,18].

Although both USG and fiberoptic evaluation were based on the evaluation of the anatomic relationship between the i-gel noninflating cuff and laryngeal entrance, there were some inequalities in our results. In our study, the specificity was 91.84% with an NPV of 91.84% to detect an acceptable i-gel placement according to the FOL grade of USG, regarding the diagnostic performance of USG in terms of FOL and LT. The specificity was 100% with an NPV of 82.65% to detect an acceptable i-gel placement based on the LT grade of USG. In other words, USG has high concordance with FOL and LT when the i-gel position is optimal. However, USG was found to have a sensitivity of 20%, with a positive predictive value (PPV) of 20% to detect unacceptable i-gel placement according to FOL grade. This result, on the other hand, shows that USG has a poor concordance in detecting i-gel malposition when compared to FOL and LT. The reason for this may be that, especially in young children, even if ventilation is sufficient, it may give an unacceptable FOL appearance, although it gives an acceptable appearance on USG due to its more anterior and highly located larynx, floppy epiglottis, larger tongues, small-sized laryngeal, and oropharyngeal anatomy [19].

Children are known to be more susceptible to complications than adults from the use of SADs [20]. The most common complications are minor and transient and include sore throat, difficulty swallowing, and hoarseness. Serious complications are very scarce and include oropharyngeal trauma, nerve damage or stomach contents aspiration [21]. In our study, broncho/laryngospasm and bradycardia were observed in only one patient (0.9%). The results were consistent with the literature [21,22].

<table>
<thead>
<tr>
<th>Test results</th>
<th>n = 108</th>
</tr>
</thead>
<tbody>
<tr>
<td>FOL image (1/2/3/4/5), n (%)</td>
<td>63 (58.3) / 28 (25.9) / 7 (6.5) / 7 (6.5) / 3 (2.8)</td>
</tr>
<tr>
<td>FOL grade, n (%)</td>
<td>98 (90.7) / 10 (9.3)</td>
</tr>
<tr>
<td>THT (0/1/2), n (%)</td>
<td>86 (79.6) / 17 (15.7) / 5 (4.6)</td>
</tr>
<tr>
<td>TLS (0/1), n (%)</td>
<td>102 (94.4) / 6 (5.6)</td>
</tr>
<tr>
<td>PPL (0/1), n (%)</td>
<td>106 (98.1) / 2 (1.9)</td>
</tr>
<tr>
<td>USG score (0/1/2/3), n (%)</td>
<td>86 (79.6) / 12 (11.1) / 7 (6.5) / 3 (2.8)</td>
</tr>
<tr>
<td>USG grade, n (%)</td>
<td>98 (90.7) / 10 (9.3)</td>
</tr>
<tr>
<td>Leakage test score, mean ± SD (min – max)</td>
<td>24.51 ± 6.40 (13–39)</td>
</tr>
<tr>
<td>Leakage test grade, n (%)</td>
<td>81 (75) / 27 (25)</td>
</tr>
</tbody>
</table>

Data are given as mean ± SD (min – max) or number (%). FOL: fiberoptic laryngoscopy, THT: transverse plane between hyoid bone and thyroid bone, TLS: transverse plane of lateral suprasternal notch, PPL: parasagittal plane of pharynx and larynx, USG: ultrasonography.
The strength of the study was the employment of separate anesthesiologists for anesthesia induction, FOL, USG imaging, and LT to overcome observer bias. In order to avoid observer bias in the USG imaging, the anesthesiologist was trained in airway imaging of SADs for about 1 week by the radiology department. Another is that we used Song et al.’s system to rule out observer bias and obtain a standard view in a performance-dependent and subjective assessment [23] such as in a USG examination. Nevertheless, there were some limitations in our study. First, we used i-gel, a second-generation SGA with a silicone body and noninflatable cuffs, but we cannot generalize our results to other structurally different second-generation SGAs as the degrees of intraoral sealing may be affected.

### 5. Conclusion

Ultrasound examination of the airway is strongly correlated with FOL and LT grade when determining the position of an i-gel and is highly compatible with acceptable positioning. USG has demonstrated very good diagnostic performance in the acceptable placement of i-gels according to both FOL and LT. However, USG showed poor diagnostic performance in confirming malpositioned i-gels.

**Table 4. Frequency table and agreement statistics of LT grade, USG, and FOL grade.**

<table>
<thead>
<tr>
<th></th>
<th>FOL grade</th>
<th>Leakage test grade</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FOL-A</td>
<td>FOL-U</td>
</tr>
<tr>
<td>USG grade, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>USG-A</td>
<td>90 (91.8)</td>
<td>8 (80)</td>
</tr>
<tr>
<td>USG-U</td>
<td>8 (8.2)</td>
<td>2 (20)</td>
</tr>
<tr>
<td>Total (%)</td>
<td>98 (90.7)</td>
<td>10 (9.3)</td>
</tr>
</tbody>
</table>

McNemar test, p-value

<table>
<thead>
<tr>
<th>Agreement statistics</th>
<th>χ², p-value</th>
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<tbody>
<tr>
<td>κ ± SE</td>
<td>0.118 ± 0.096, p = 0.218, Z = 1.23, p = 0.468 ± 0.081, Z = 5.75, p &lt; 0.001</td>
</tr>
<tr>
<td>Proportion A agreement</td>
<td>91.84%</td>
</tr>
<tr>
<td>Proportion U agreement</td>
<td>20%</td>
</tr>
<tr>
<td>Diagnostic measures (% 95% CI) for unacceptable Sensitivity Specificity PPV</td>
<td>20 (5.25–55.61), 91.84 (84.55–96.41)</td>
</tr>
<tr>
<td>Accuracy</td>
<td>85.19 (77.06–91.29)</td>
</tr>
<tr>
<td>Incidence rate for U in USG (% 95% CI) (9.26 % (17.17–34.25)</td>
<td></td>
</tr>
<tr>
<td>Incidence rate for A</td>
<td>83.33% (74.94–89.81)</td>
</tr>
<tr>
<td>Incidence rate for U</td>
<td>9.26 % (17.17–34.25)</td>
</tr>
</tbody>
</table>

Comparison of incidence rate for U with Z-test with χ² test

<table>
<thead>
<tr>
<th>Relationship between methods (for numerical data)</th>
<th>p &gt; 0.999</th>
<th>p &gt; 0.999</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spearman’s rho</td>
<td>r = 0.203, p = 0.035</td>
<td></td>
</tr>
<tr>
<td>Kendall’s τ B</td>
<td>r = 0.191, p = 0.032</td>
<td></td>
</tr>
<tr>
<td>r = 0.345, p &lt; 0.001</td>
<td>r = 0.345, p &lt; 0.001</td>
<td></td>
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</tbody>
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**Informed consent**

Ethics committee approval [Selçuk University Faculty of Medicine the Local Ethics Committee (14.10.2020.2020/19)] and Clinical Trials Registry (ClinicalTrials.gov Identifier: NCT04652466) was obtained for this study and all participants signed an
informed consent form.
References


