Comparing S-Guide® and Gliderite® Times to Assist Video Laryngoscopic Intubation in Patients with Simulated Difficult Airways: A Single-Blinded Randomized Prospective Study

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Abstract

Objective: Gliderite®, one of the first stylets designed specifically to assist tracheal intubation with non-channeled curved blades video laryngoscopes, can cause injury. The S-Guide® is a new, malleable, intubating guide allowing oxygenation. Its soft tip is designed to prevent trauma. We aimed to compare the duration of tracheal intubation with S-Guide compared to Gliderite using a C-MAC® D-Blade® video laryngoscope in patients with simulated difficult airways.

Methods: We performed a single-blinded prospective randomized study, with 50 adult patients requiring orotracheal intubation under general anaesthesia in Lausanne University Hospital. A cervical collar was fitted around the patient’s neck to simulate difficult intubation conditions. Exclusion criteria were American Society of Anesthesiologists (ASA) >3, BMI >35 kg m2, known or at risk of difficult intubation, and risk of aspiration of gastric content. We recorded T1: time of identification of the glottis; T2: time to inflate the cuff, and T3: total intubation time (capnography curve appearance). Secondary outcomes were the presence of arytenoid contact during intubation and postoperative airway discomfort.

Results: There were no significant differences between T1 and T2 (seconds) while using the S-Guide or Gliderite, respectively: 14.6 [9.6-18.6] vs 16.5 [11.0-20.6]; P = .368 and 43.3 [33.2-49.3] vs 46.3 [35.6-61.5], P = .308. T3 was significantly shorter in the S-Guide group: 58.1 [50.2-61.8] vs 65.3 [57.6-78.7], P = .044. Fewer arytenoid contact occurred during intubation using the S-Guide (P = .032), without difference in postoperative airway discomfort.

Conclusion: S-Guide-assisted tracheal intubation, with a C-MAC D-Blade in simulated difficult airways, allows successful and faster intubation than with the Gliderite Stylet.

Keywords: Difficult airways, intubation trauma, stylet, video laryngoscopic intubation

Main Points

- The S-Guide is a new malleable, intubating stylet, with a soft colored tip, designed to prevent trauma and allow oxygenation through its hollow lumen.
- In a single-blinded prospective randomized study including 50 adults, the S-Guide-assisted tracheal intubation with a C-MAC D-Blade, in simulated difficult airways, allowed successful and faster intubation.
- The comprehensive uses of the newly designed S-Guide for intubation will need further investigation.
Introduction

Tracheal tube introducers (bougie) and stylets are essential tools in difficult airway management\(^1\)\(^2\)\(^3\)\(^4\)\(^5\)\(^6\)\(^7\)\(^8\)\(^9\)\(^10\) with reported success rates from 78\%\(^{11}\)\(^\text{--}\)100\%.\(^{12}\) Various complications related to their extended use range from mild sore throat to mucosal bleeding and bronchial or palatopharyngeal perforation.\(^{9}\)\(^{11}\)\(^{12}\)\(^{13}\)\(^{14}\)

Although first described in 1949 by Sir Robert Macintosh\(^{10}\), while using a urethral catheter (hence the popular term of Gum Elastic Bougie or GEB), improvements in manufacturing, technology and understanding airways have resulted in numerous modifications of the original device.\(^1\)\(^2\)\(^3\)\(^4\)\(^5\)\(^6\)\(^7\)\(^8\)\(^9\)\(^10\)\(^11\)\(^12\)\(^13\)\(^14\)\(^15\)\(^16\)\(^17\)\(^18\)\(^19\)\(^20\)\(^21\)\(^22\)\(^23\)\(^24\)\(^25\)\(^26\) The manufacturer of the GlideScope\(^{\text{®}}\) has specifically designed a reusable stylet for VLS.\(^{18}\) The Gliderite\(^{\text{®}}\) Rigid Stylet (Verathon Medical Europe B.V. Amsterdam, The Netherlands: GlideScope video intubation system operator and service manual) is reusable and more rigid than standard malleable stylets. Its length of 266 mm and outer diameter allow railroading a tube of size 6 (ID) mm and greater. The distal curvature approaches 90°\(^\text{°}\) and its radius of curvature is approximately 6 cm. It does not allow oxygenation.\(^{9}\) The potential for injury has however been highlighted, despite its specific design.\(^{20}\)\(^{21}\)\(^{22}\)\(^{23}\)

The 15 French (Fr) S-Guide\(^{\text{®}}\) is a new single-use, flexible, multifunctional intubating guide (VBM Medizintechnik GmbH, Einsteinstrasse 1, D-72172 Sulz a.N.). Its color-coded soft tip is designed to prevent trauma during intubation (see Figure 1 and Figure 2).\(^{21}\)\(^{22}\) Its metallic core allows malleability with shape retention\(^{25}\) and oxygenation is possible through its hollow lumen.\(^{21}\)\(^{22}\) Recent developments have allowed 2 new sizes to be produced (11 Fr and 8 Fr), allowing railroading of tubes sized 4.5 and 3.0, respectively.

To the best of our knowledge, there is no available evidence comparing the performance of the 15-Fr sized intubating guide with an established intubation stylet to assist VLS intubation.

We hypothesized that, in a simulated difficult airway setting, the total time for intubation using a C-MAC D-Blade would be significantly shorter if the intubation procedure was assisted with the S-Guide stylet instead of the Gliderite. Through observation of the intubation technique, we also took interest in postoperative throat discomfort and tried to see if there was any correlation with arytenoid contact.

We aimed to conduct a single-blinded randomized controlled trial to compare the S-Guide intubating guide with the specifically designed Gliderite\(^{\text{®}}\) stylet to assist C-MAC\(^{\text{®}}\) D-blade\(^{\text{®}}\) video laryngoscopic tracheal intubation for patients with simulated difficult airways.

Methods

All procedures performed in this study involving human participants were in accordance with the ethical standards of the Human Research Ethics Committee of the Canton Vaud (July 7, 2015, protocol 267/15, Chairperson Prof. Patrick Francioli) and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. Written informed consent was obtained from all subjects participating in the study. The study was registered prior to patient enrollment at www.clinicaltrials.gov (NCT02519647, Principal investigator: Schoettker Patrick, Date of registration: August 11, 2015). This prospective, patient-blinded, randomized controlled trial was designed to compare time necessary to intubate, success rates, ease of intubation, and postoperative complications due to tracheal intubation assisted by the S-Guide or the Gliderite using the C-MAC D-blade in patients with a difficult airway simulated by a cervical collar.\(^{11}\)\(^{26}\) We included 50 adult patients, with ASA physical status 1-3, scheduled for elective surgery at Lausanne University Hospital and requiring orotracheal intubation under general anaesthesia (Figure 1). Exclusion criteria were patients with a BMI > 35 kg m\(^2\), known difficult intubation, an interdental distance < 3.5 cm, a thyromental distance < 6 cm, or at risk of aspiration of gastric content. Patient recruitment and follow-up took place from August 1, 2015, until June 2, 2016.

The anaesthesia protocol has been published previously.\(^{26}\)\(^{27}\) A cervical collar was fitted around patients’ necks before intubation in order to reduce mouth opening and limit cervical movement in order to simulate a difficult airway, as already published in a previous study.\(^{26}\) Patients were randomly assigned to the Gliderite or S-Guide group using a computer-generated randomization list (www.randomization.com). The disclosure of the intubating device was done by the supervisor just before the beginning of each procedure. Both devices were available as part of our department’s equipment and both devices were bought at market price without any fundings. Gliderite and S-Guide were available as part of our department’s equipment and were bought at market price.

All tracheal intubations were performed using a standard 6.5 mm cuffed tube for women patients and a 7.5 mm tube for men (Mallinckrodt\(^{\text{®}}\) Hi-Contour Oral Tracheal Tube...
Cuffed; Covidien llc, 15 Hampshire Street, Mansfield, Mass, USA). The Gliderite and S-Guide were lubricated with silicon spray before insertion into the tube and used according to manufacturers’ instructions.

The timer was started on contact with the C-MAC (T0). Time of identification of the glottis was recorded as T1 (expressed as median seconds [25th-75th]), and time of blocking the cuff was recorded as T2. Time of ventilation was defined as the time needed to see the end-expiratory CO2 curve on capnography and also represented the total intubation time T3 (time to CO2) as defined in our previous protocol.26

Success and the number of attempts necessary were recorded. Tracheal intubation was considered as failed if it could not be accomplished within 3 minutes or in the event of desaturation (SpO2 < 92%).

All patients’ tracheas were intubated, under the first author’s supervision, by trainee anaesthetists to avoid a potential bias toward any specific equipment that senior anaesthetists could have. All had at least 1 year of experience in anaesthesiology and had used the C-MAC D-blade more than 5 times previously in a clinical setting. Both the intubating doctor and the supervisor assessed the subjective ease of the intubation procedure on a scale from 1 (very easy) to 5 (very difficult). Ease of insertion of the D-blade, ease of glottis identification, and ease of insertion of the tracheal tube through the tracheal inlet were also assessed. The supervisor also recorded whether there was contact with the arytenoid during the intubation process.

Postoperative discomfort was assessed 24 hours after intubation, identifying the presence of a sore throat (pain score from 1 to 5), hoarseness, dry throat, or dysphagia.

The primary outcome was the total intubation time. Secondary endpoints included successful intubation and number of attempts necessary, the times of glottis identification, inflating the cuff, and apparition of end-expiratory
CO₂, as well as the subjective ease of intubation and postoperative discomfort.

Sixteen different anaesthetists took part in the study, each performing from 1 to 8 tracheal intubations, reflecting the clinical setting of a teaching hospital center. At the end of the study, the anaesthetists involved were asked to rate the devices they had used and give 1 positive and/or 1 negative comment about the device of their choice.

Statistical Analysis

Based on a reference established by Bathory et al.26 in a similar model of difficult intubation, we identified a 20% shorter intubation time for the S-Guide group to be clinically relevant. Sample size calculation yielded a required sample size of n = 25 per group to detect statistically significant group differences with an α error of 0.05 and a power of 80%.

All statistical analyses were performed using Stata software (v. 14.2, StataCorp, College Station, Tex, USA). Categorical data are presented as raw frequencies and relative percentages. Distribution differences in the categorical data between 2 or more independent groups were assessed using the chi-square test or Fisher’s exact test in cases of insufficient sample size. Distributions of continuous data were first evaluated using normal QQ-plots. Gaussian distributed data were summarized as mean, standard deviation, and range, whereas non-Gaussian distributed data were summarized as median, interquartile range, and range. Differences in means between 2 independent groups of Gaussian distributed data were assessed using Student’s t-test; for non-Gaussian distributed data, the non-parametric Mann–Whitney–Wilcoxon

Table 1. Characteristics of Patients Randomly Assigned to Gliderite or S-Guide

|                              | Gliderite (n = 25) | S-Guide (n = 25) | P  |
|------------------------------|--------------------|------------------|----|
| Sex                          |                    |                  |    |
| Female                       | 11 (44.0)          | 9 (36.0)         | .564|
| Male                         | 14 (56.0)          | 16 (64.0)        |    |
| Weight in kg (median, mean, (SD), IQR [range]) | 72, 75.3, (16.8) 65-82 [50-128] | 74, 73.7, (14.3) 62-85 [48-99] | .961|
| Height in cm (median, mean, (SD), IQR [range]) | 172, 170.9, (9.6) 164-178 [154-186] | 170, 169.1, (9.7) 165-175 [148-185] | .586|
| BMI in kg/m² (median, mean, (SD), IQR [range]) | 24.7, 25.7, (5.0) 22.0-27.5 [19.8-39.1] | 24.5, 25.7 (4.9)21.8-29.4 [19.3-36.2] | .977|
| Age in years (median, mean, (SD), IQR [range]) | 55, 55.3, (15.1) 46-66 [23-92] | 61, 58.2, (15.8) 49-66 [28-91] | .610|
| ASA status                   |                    |                  |    |
| I                            | 1 (4.0)            | 0 (0.0)          | 1.000|
| II                           | 19 (76.0)          | 20 (80.0)        |    |
| III                          | 5 (20.0)           | 5 (20.0)         |    |
| Mallampati score             |                    |                  |    |
| 1                            | 9 (36.0)           | 7 (28.0)         |    |
| 2                            | 14 (56.0)          | 15 (60.0)        |    |
| 3                            | 2 (8.0)            | 2 (8.0)          |    |
| 4                            | 0 (0.0)            | 1 (4.0)          | .905|
| TMD in cm (median, mean, (SD), IQR [range]) | 7.5, 7.6, (0.6) 7.0-8.0 [6.5-9.0] | 7.5, 7.4, (0.5) 7.0-8.0 [6.5-8.5] | .466|
| IDD in cm (median, mean, (SD), IQR [range]) | 4.2, 4.0, (0.7) 3.5-4.5 [2.4-5.8] | 4.2, 4.2, (0.6) 3.9-4.5 [2.8-5.6] | .315|

Data are shown as number [percentage], mean value, median, standard deviation (SD), and interquartile range (IQR) [range].

TMD, thyromental distance; IDD, interdental distance; BMI, body mass index.
A rank-sum test was used. A $P < .05$ was considered statistically significant. The presence of statistically significant differences of co-founding factors between the 2 groups in terms of ASA status, weight, height, and factors predictive of difficult intubation were tested also tested through Student’s $t$-tests and Mann–Whitney–Wilcoxon tests.

## Results

Fifty patients were randomly attributed to 2 groups without any statistically significant differences noted in terms of sex, ASA status, weight, height, and factors predictive of difficult intubation (Table 1).

All the patients’ trachea were intubated successfully except for 1 patient in the Gliderite group. For this particular individual, tracheal intubation was eventually successful using the S-Guide as a rescue tool. None underwent desaturation.

No significant differences were measured in times of glottis identification $T_1$ (second): 14.6 [9.6-18.6] vs 16.5 [11.0-20.6]; $P = .368$ or cuff blocking $T_2$ (second) 43.3 [33.2-49.3] vs 46.3 [35.6-61.5]; $P = .308$, for the S-Guide and Gliderite groups, respectively (Figure 2). The total intubation time (time to CO$_2$), $T_3$ (second) was significantly shorter in the S-Guide group: 58.1 [50.2-61.8] vs 65.3 [57.6-78.7]; $P = .044$.

Concerning our secondary endpoints, the trainee anaesthetists and supervisor subjectively considered the tracheal intubation to be significantly easier with the S-Guide (Table 2). There were no differences between the 2 groups with regards to D-blade insertion difficulty or glottis identification.

Significantly less contact with the arytenoids was observed with the S-Guide (13 vs 20; $P = .032$). Postoperatively, S-Guide group patients experienced overall less discomfort yet not significantly, as no significant correlation was established (Table 3).

Decreased trends for each individual variable assessed were reported (Table 4). One patient in the Gliderite group underwent Tracheotomia. The presence of a hoarse voice couldn’t be evaluated.
Overall, anaesthetists favored usage of the S-Guide (9 rated the S-Guide higher than or at least equal to the Gliderite; 6 only used one of the devices and therefore could not be compared; 1 rated the Gliderite higher).

Negative comments concerning the S-Guide included the potential need for a “3-handed intubation procedure,” with the third hand mainly needed to withdraw the S-Guide to allow for tube movement (2 negative comments). Two anaesthetists made negative comments about the soft-tipped curved end and 2 complained about an involuntary rotation movement of the S-Guide within the tube.

Negative comments about the Gliderite mainly concerned difficulties in positioning the tube between the vocal cords and the need to sometimes forcefully withdraw the device after tracheal intubation.

Discussion

Recent studies highlighted that differences in tracheal intubation times were dependent on devices and operators, especially when using VLS technology instead of Macintosh intubation.

This randomized controlled trial shows that the success of tracheal intubation performed with a C-MAC D-Blade in patients with a simulated difficult airway was not significantly different between the use of a 15 Fr S-guide and a Gliderite. Although tracheal intubation times were significantly shorter in the S-guide group, we did not demonstrate the 20% time reduction of the total time of intubation (T3), initially expected while designing the present study.

Tracheal intubation for VLS requires tube handling and positioning to allow delivery through the tracheal inlet. While anatomical visualization using unchanneled VLS can generally be described as good, the success rate of tracheal intubation increases with the usage of a stylet or bougie. Various authors have described specific distal curvatures, ranging from 60° to 90°, but no specific curve has shown overall superiority. The S-Guide was used according to the manufacturer’s instructions and bent by the user into a hockey stick shape, which is slightly less angulated than the Gliderite.

Subjectively, the color-coded soft tip of the S-Guide allowed anaesthetists easier positioning in front of the tracheal inlet, better aim and positioning between the vocal cords, and streamlining the process of tracheal intubation. A similar technique could not be achieved with the Gliderite, which could furthermore lead to potential airway trauma, due to its rigidity and hard tip.

Stylet-assisted tracheal intubation for VLS has been described as responsible for airway trauma. Our study revealed no significant differences in postoperative airway discomfort with regards to sore throat, throat pain score, hoarse voice, or dysphagia. However, every single item showed a diminished incidence in the S-Guide group. Less arytenoid contact was described in the use of the S-Guide, whereas no significant correlation could be established between arytenoid contact and postoperative discomfort. A soft-tipped bougie and associated lower arytenoid contact might be independent characteristics contributing to a decrease in postoperative airway discomfort, although this has variable origins, ranging from mucosal lacerations to arytenoid dislocation. The present study documented no clinically relevant injuries, and all tracheal intubations were performed safely.

The present study has some limitations. First, our study simulated difficult airway management by using a semi-rigid collar, limiting mouth opening and neck extension. It did not assess the S-Guide’s performance in comparison with the Gliderite in a variety of difficult intubation scenarios. In cases involving
airway malignancies or disrupted anatomy, the performances of both devices might differ from our results, and this needs further assessment.

Second, although we were able to show a statistically significant time reduction in the S-Guide group, a difference of 7 seconds might not be clinically relevant. We however believe that a reduction of more than 10% of the total intubation time contributes to better airway management in patients with simulated difficult airways.

Third, no significant differences in postoperative airway discomfort were revealed in this study. Yet, our study was not powered to assess potential outcome on throat injury and the sample size was relatively small. New studies should be carried out in different clinical cohorts, especially with the newly sized pediatric 11 Fr and neonatal 8 Fr. S-Guide have been made available on the market. Follow-up multi-center study is necessary to generalize the conclusion of this study.

Furthermore, even though oxygenation is possible through its hollow lumen, no patient presented episodes of desaturation in any group. Further investigations are necessary to assess the clinical significance of this option.

Fourth, the Gliderite stylet was originally designed to assist intubation using the GlideScope® VLS. Yet, in our study, we used a single video laryngoscope model which was the C-MAC with D-Blade. This might be seen as a potential bias as the curvature differs between the 2 set ups.

Finally, although every effort was taken to minimize any conflicts of interest, the present study’s senior author was part of the S-Guide’s design team. This might have influenced results in terms of a bias in the intubating anaesthetists’ responses. However, all procedures were performed by trainee anaesthetists, thus reducing the risk of any consolidated preference for any specific intubation system. In addition, the senior anaesthetist was neither present in the operating theatre when intubation was performed nor was he involved in data collection.

**Conclusion**

The use of the newly designed S-Guide compared to the Gliderite for successful intubation will need further investigation. The S-Guide stylet can be seen as a new helpful tool in the management of the difficult airway available to the anaesthetist, intensivist, or the emergency physician. Its single-use profile can be seen as an advantage in pandemic situation.

Recent case reports have shown the S-Guide utility either in an out-of-hospital emergency settings or in a situation of subglottic stenosis. Its color-coded soft tip is considered as an advantage to ease its precise positioning between the vocal cords without fearing of hurting patients and its malleability might help to overcome anatomic barriers in the oropharyngeal tract.

This study did not assess the possibility of oxygen delivery through the S-Guide. While this option is a promising tool for patients with low oxygen reserve, its usefulness is also expected in situations where the intubation procedure is feared to be time-consuming.

These clinically relevant advantages represent an opportunity for further research.

Based on the present findings, our department has added the S-Guide to its range of primary learning tools for dealing with difficult airway, especially for young trainees who are less experienced. Emphasis on understanding, teaching, and training has further been implemented.

**Ethics Committee Approval:** The study was approved by the Human Research Ethics Committee of the Canton Vaud (July 7th, 2015, protocol 267/15, Chairperson Prof. Patrick Francioli). Clinical trial number and registry URL: NCT02519647 www.clinicaltrials.gov.

**Informed Consent:** Written informed consent was obtained from all subjects participating in the study.

**Peer-review:** Externally peer-reviewed.

**Author Contributions:** Concept – C.N., T.M., I.B., P.S.; Design – C.N., T.M., I.B., P.S.; Supervision – P.S.; Resources – P.S.; Materials – P.S.; Data Collection and/or Processing – C.N., T.M., I.B., P.S., N.F.; Analysis and/or Interpretation – C.N., T.M., I.B., P.S., N.F.; Literature Search – C.N., T.M., I.B., P.S.; Writing Manuscript – C.N., T.M., P.S.; Critical Review – C.N., T.M., I.B., P.S., N.F.

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**Declaration of Interests:** The present study’s senior author was part of the S-Guide’s design team. However, he did not participate in any intubations or data collection himself.

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