INTRODUCTION

Endotracheal intubation is a definitive way of securing the airway and is routinely done by laryngoscopy and visualisation of cords. However, this involves distortion of upper airway to bring glottis into the line of sight\(^1\) and in some situations such as high larynx, facial trauma, etc., tracheal intubation fails. Supraglottic airway devices (SADs) are useful in such situations for rescue ventilation. Laryngeal mask airway (LMA) classic (c-LMA)\(^2\) is one such device which is included in Difficult Airway Society guidelines for unanticipated difficult intubation.\(^3\)

Laryngeal mask airway classic was designed for maintenance of airway in emergency situations, especially by untrained personnel. Later it was modified into intubating LMA (ILMA) or LMA Fastrach\(^4\). Major difference between standard LMA and LMA Fastrach\(^5\) lies in the design and function of the shaft which is rigid as compared to soft silicone shaft of c-LMA thus facilitating adjusting

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manoeuvres to align the mask’s aperture against the glottis opening.

The i-gel® is a relatively new single-use SAD which does not have an inflatable cuff.[4] It is made from a soft, gel-like and transparent thermoplastic elastomer (styrene ethylene butadiene styrene) which creates a noninflatable seal which is a mirror impression of the supraglottic anatomy.[5] The i-gel® has several other useful design features including a gastric channel, an epiglottic ridge and a ridged flattened stem to aid insertion and reduce the risk of axial rotation.[6] The stem of the i-gel® is less flexible than that of the LMA-classic and has an integral bite.[7] i-gel® has also been used in rescue airway management and as a conduit for tracheal intubation.[6-12]

The aim of our study was to compare the success rate of blind tracheal intubation through the i-gel versus the LMA Fastrach®. Because of higher airway leak pressure[13-15] and better visualization of glottis,[16] as compared to LMA Fastrach®, we assumed a better first-attempt success rate during blind tracheal intubation through i-gel®.

**METHODS**

After receiving approval from Ethical Committee and informed consent from patients, 100 patients were randomly assigned using a chit method into two groups of 50 each: The LMA Fastrach group (group F) and the i-gel group (group I). Patients between 18 and 60 years of age, American Society of Anaesthesiologists (ASA) grade I/II, with adequate mouth opening who required general anaesthesia and tracheal intubation for an elective surgical procedure were included in the study. Exclusion criteria were ASA class > II, mouth opening < 2 cm, increased risk of aspiration, oral pathology and known or anticipated difficult tracheal intubation or facemask ventilation. Injection ranitidine 50 mg and metoclopramide 10 mg were administered intravenously 30 min before the operation. In the operation theatre, infusion of Ringer lactate solution was started. Patients were connected to a multichannel monitor showing electrocardiography, oxygen saturation, noninvasive blood pressure and end-tidal carbon dioxide (EtCO₂). All patients were administered injection glycopyrrolate 0.2 mg, injection midazolam 0.02 mg/kg and injection fentanyl 2 µg/kg IV. Preoxygenation was done for 3 min. All patients were induced with injection propofol 2 mg/kg IV in slow incremental dose and ease of mask ventilation was noted. After confirming adequate mask ventilation, injection rocuronium 0.9 mg/kg was administered to facilitate intubation. The supraglottic device was inserted once the jaw relaxation was achieved. The investigator had already gained experience of blind intubation in 50 patients with LMA Fastrach® and i-gel® each before the study. These cases were not included in the study.

Selection of size of the LMA Fastrach® and the i-gel® was on the basis of the weight of the patient. For the lubrication of the SADs, we used water-based lubricating jelly. Both SADs were introduced as per manufacturer's user booklet.[4,17] The i-gel® was inserted in sniffing morning air position, while the LMA Fastrach® was inserted in neutral neck position. Adequate ventilation was confirmed by chest movements and EtCO₂ waveforms. When ventilation was not adequate, or there was an audible leak, while ventilating with an inspiratory pressure of 20 cm H₂O, up down manoeuvre causing change in depth of insertion was performed. Use of different size of SAD were attempted if required.

Time required for insertion of SAD was defined from removal of the facemask to the time where adequate ventilation was established through SAD with capnographic confirmation.

Conventional polyvinyl chloride (PVC) endotracheal tubes (ETTs) were used for blind tracheal intubation in both the groups. ETT was lubricated with water-based lubricating jelly prior to insertion. Size 7.0 mm internal diameter (ID) ETT was used in patients weighing > 50 kg and 6.0 mm ID ETT for patients < 50 kg.[12]

In group F, ETT was inserted with reverse orientation.[10-20] In this manoeuvre, ETT is rotated through 180° once it crosses the proximal opening in LMA. If resistance was encountered during insertion of tracheal tube in F group, a standardised algorithm was followed on the basis of the distance at which the resistance was felt, as recommended by manufacturer.[21] If no resistance was felt during insertion of tracheal tube, it was advanced fully into the ILMA. Intubation was considered successful, if adequate ventilation produced a chest rise and a capnographic waveform.

In group I, ETT was rotated 90° counter-clockwise during insertion [Figures 1 and 2].[12] This method was used by Halwagi et al. and resulted in increased success rate, so it was incorporated in our study. If the resistance was felt during insertion, i-gel® was
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readjusted and stabilised at the point of maximum chest expansion, and an assistant was asked to perform optimum external laryngeal manipulation (OELM) by applying backward pressure on thyroid cartilage. This resulted in increased success rate of intubation and prevents the impingement of bevel on glottis structures. However, the disadvantage was that we had to take the help from the second person to perform external laryngeal manipulation, but the results were satisfactory.

Time required for blind endotracheal intubation was defined from disconnection of the breathing circuit of the SAD to confirmation of the tracheal intubation by auscultation and capnograph trace.

After confirmation of correct placement of ETT, supraglottic device was removed using a stabilizing rod.

In both study groups, two attempts at device insertion and intubation were allowed. Intubation was only attempted if appropriate ventilation was obtained through SAD. If tracheal intubation through the device was unsuccessful, it was performed by direct laryngoscopy.

During intraoperative and postoperative period and upto 24 h patients were evaluated for any adverse event or postoperative complaints such as sore throat, pain on swallowing and hoarseness.

The sample size was calculated to detect a 10% difference in first-attempt success rate in ETT insertion between devices with a type-I error of 0.05 and a power of 90%, requiring 25 patients per group. We included 50 patients in each group to allow for potential drop-outs. Normally distributed data were analysed using t-test, and categorical data were analysed using the Chi-square test. Continuous data are presented as mean and standard deviation, whereas categorical data are presented as number of patients and percentage. Data were analysed using IBM SPSS statistics 20.0 software. $P < 0.05$ was considered statistically significant.

**RESULTS**

Out of 147 patients that were evaluated for the study, 41 patients were excluded according to the exclusion criteria. Six patients did not give informed consent, hence, we randomized 100 patients in two groups of 50 each. Demographic data were similar in both the groups [Table 1].

There was no difference in the successful insertion of SADs between the two groups, that is, i-gel and ILMA. With the first attempt of SAD insertion, the successful ventilation rate was 96% in I group and 90% in F group. With the second attempt of SAD insertion, the successful ventilation rate was 100% in both the groups [Table 2]. With the first attempt, blind tracheal intubation was successful in 66% cases (33 patients) of I group and in 74% cases (37 patients) of group F. With the second attempt, blind tracheal intubation was successful in 82% cases (41 patients) of group I and 96% cases (48 patients) of group F [Table 2]. Total time to achieve successful ventilation with SAD was shorter in group I. Time for successful ventilation with SAD was 19.40 s in the group I and 38.96 s in group F ($P < 0.0001$). Time to achieve successful intubation through the SADs was 24.04 s in the group I when compared to 20.96 s in group F ($P = 0.103$) [Table 2].
Postoperative complications in both the groups were comparable. However, dysphonia was more in the group I but still i-gel proved to be slightly safer than LMA Fastrach [Table 3].

**DISCUSSION**

In this study, overall success rate of insertion of supraglottic devices in both the groups was 100% which was similar to various previously conducted studies. In the present study, first-attempt success rate for blind tracheal intubation was comparable in both the groups and overall success rate was higher in F group as compared to I group, which is similar to the results of Halwagi et al. (2012) and Sastre et al. (2012) who noticed higher success rate of blind tracheal intubation with ILMA. This could be due to a “V” shaped tracheal tube guiding ramp in LMA Fastrach that centralizes the ETT towards the glottic aperture as the ETT emerges from the metal shaft and guides it anteriorly to reduce the risk of arytenoids trauma and oesophageal placement and the presence of the handle in LMA Fastrach which resulted in stabilization and manipulations which could not be done in i-gel. So in group I, we did external manipulation of the larynx. In our study, when first attempt of blind intubation was unsuccessful in group I, we stabilized the i-gel at the point of maximum chest expansion by readjusting and took the help of an assistant to apply external laryngeal pressure. This resulted in better overall success rate of ETT insertion through i-gel (82%) as compared to studies by Halwagi et al. (73%) and Sastre et al. (40%). In group F ETT was inserted with reverse orientation as this method resulted in higher success rate in various studies. It optimises the ETT with the angle of trachea resulting in better first-attempt success rate of ETT insertion.

The overall intubation success rate using LMA Fastrach was comparable to published studies. The cases in which blind tracheal intubation failed only two patients needed stylet for intubation with Macintosh laryngoscope in group I and none in group F. The easier and a quicker insertion of i-gel was probably due to noninflation of cuff. Time was not wasted in inflating the cuff, and moreover, the rigid structure of LMA Fastrach causes delay in insertion as compared to i-gel.

i-gel has been used as a conduit for tracheal intubation with the help of fibreoptic bronchoscope in several case reports. But limited studies are there to prove the efficacy of use of i-gel for blind intubation. Michalek et al. did blind tracheal intubation in three different airway manikins through the i-gel with a success rate of 51%. Theiler et al. studied “visualised blind intubation” through the i-gel and the LMA Fastrach. Their results showed a poor success rate (15%) with i-gel as compared with the LMA Fastrach (69%). Sastre et al. also showed an inferior intubation rate of 40% through i-gel as compared to 70% with LMA Fastrach. We observed that 90° counter-clock rotation and OELM resulted in substantially superior results.

The incidence of postoperative complications was comparable in both the groups. In the present study,
dysphonia was more in I group which was similar to study conducted by Sastre et al. While the incidence of sore throat was lesser in I group when compared to F group; this observation is similar to that of Keijzer et al.\[15\]

There are however some limitations to our study. Data were collected in an unblended manner, some bias was possible. All the patients were ASA grade I or II with no anticipated difficult intubation. This does not represent the general population. Furthermore, the investigator had less prior experience with i-gel\[8\] as compared to LMA Fastrach\[7\].

The use of wire reinforced tubes would have resulted in better success rate, but conventional PVC tubes are readily available and cheaper. However, in LMA Fastrach\[7\], there was no difference in successful blind tracheal intubation with conventional tracheal tube and silicon wire-reinforced tracheal tube in studies conducted by Lu et al.\[19\] and Kundra et al.\[26\] but in case of i-gel further studies are required.

**CONCLUSION**

We concluded that i-gel\[8\] can be used as a conduit for endotracheal intubation. Though it is an effective SAD, it is slightly inferior to LMA Fastrach\[7\] as the intubating device. Further studies are required to prove its efficacy as a conduit for intubation.

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Announcement

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