The clinical effectiveness of the streamlined liner of pharyngeal airway (SLIPA™) compared with the laryngeal mask airway ProSeal™ during general anesthesia

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Background: The aim of this study was to compare the streamlined liner of the pharynx airway (SLIPA), a new supraglottic airway device (SGA), with the laryngeal mask airway ProSeal™ (PLMA) during general anesthesia.

Methods: Sixty patients were randomly allocated to two groups; a PLMA group (n = 30) or a SLIPA group (n = 30). Ease of use, first insertion success rate, hemodynamic responses to insertion, ventilatory efficiency and positioning confirmed by fiberoptic bronchoscopy were assessed. Lung mechanics data were collected with side stream spirometry at 10 minutes after insertion. We also compared the incidence of blood stain, incidence and severity of postoperative sore throat and other complications.

Results: First attempt success rates were 93.3% and 73.3%, and mean insertion time was 7.3 sec and 10.5 sec in PLMA and SLIPA. There was a significant rise in all of hemodynamic response from the pre-insertion value at one minute following insertion of SLIPA. But, insertion of PLMA was no significant rise in hemodynamic response. There was no statistically significant difference in the mean maximum sealing pressure, gas leakage, lung mechanics data, gastric distension, postoperative sore throat and other complication between the two groups. Blood stain were noted on the surface of the device in 40% (n = 12) in the SLIPA vs. 6.7% (n = 2) in the PLMA.

Conclusions: The SLIPA is a useful alternative to the PLMA and have comparable efficacy and complication rates. If we acquire the skill to use, SLIPA may be considered as primary SGA devices during surgery under general anesthesia. (Korean J Anesthesiol 2010; 58: 450-457)

Key Words: Airway, Anesthesia, PLMA, SLIPA, Supraglottic airway.
Introduction

In general anesthesia, endotracheal intubation is used to secure the airway passage. When intubation using laryngoscopy is difficult or when one wants to reduce hemodynamic changes caused by it, a supraglottic airway device (SGA) may be used instead. The best known SGA is the laryngeal mask airway (LMA). Unfortunately it has a low sealing pressure. It can therefore cause inadequate ventilation and gastric distension during positive ventilation, and carries a risk of pulmonary aspiration in the event of regurgitation [1]. Many devices have been developed to minimize these drawbacks. One of the most commonly used SGAs is the laryngeal mask airway ProSeal™ (PLMA, Laryngeal Mask Company, UK), a modified version of the classic LMA with an additional channel for gastric tube placement to prevent pulmonary aspiration and a superior cuff that can provide airway sealing pressures higher than those of the classic LMA [1].

The newly developed SLIPA™ (stream lined liner of pharyngeal airway™, SLIPA Medical Ltd., UK) is replacing the LMA in general anesthesia. It is made of soft plastic in an anatomical shape which sticks to the pharynx and palate. It therefore does not require cuff inflation and extra straps to maintain a secure airway during positive pressure ventilation. Furthermore, a 50 ml empty internal space allows the removal of pharyngeal secretions, thus reducing the risk of pulmonary aspiration [2] (Fig. 1). Also, it can be inserted without the need for other devices, and as it is designed for single-use at a relatively cheaper price than the LMA, it has a lower risk of infection [3].

While there has been much clinical research (in Korea and elsewhere) on other SGAs, there has, especially in Korea, been little study on the SLIPA. Also, there is as of yet no comparative research on SLIPA and PLMA.

In the present study, we compared the use of the PLMA and SLIPA during general anesthesia in regard to their first insertion success rate, ease of use, hemodynamic responses to insertion, ventilatory efficiency and postoperative complications. We also evaluated the clinical effectiveness of each method.

Materials and Methods

The 60 patients chosen for the study were ASA class I–III, were between the ages 12–75 years, and were undergoing elective surgery under general anesthesia. Excluded from the study were: patients with a history of diabetes mellitus, morbid obesity, gastroesophageal reflux, and pregnancy, patients at risk of pulmonary aspiration, and patients electing for surgery in the non-supine position. After receiving the approval of the hospital’s ethics committee, all patients gave written informed consent. The patients were randomly divided into the SLIPA and PLMA groups with 30 patients in each. Randomization was based on Microsoft Excel random number generation. Patient group allocation was not revealed to investigator until immediately prior to induction of anesthesia by means of numbered, sealed envelopes. The two groups were comparable with respect to gender, age, height, weight, ASA classification, modified Mallampati classification, and the duration of anesthesia (Table 1). To evaluate the airway, the modified Mallampati classification was recorded at preanesthetic evaluation.

In the PLMA group, size 3 was used for patients weighing below 50 kg, size 4 for those weighing 50–70 kg, and size 5 for those over 70 kg. After insertion, the cuff pressure was maintained at 60 cmH₂O with a hand pressure gauge (VBM, Germany). In the female SLIPA group, size 47 was used for patients under 155 cm in height, size 49 for those between 155–165 cm, and size 51 for those over 165 cm. In the male SLIPA

Table 1. Demographics Data and Duration of Anesthesia

|                  | PLMA group (n = 30) | SLIPA group (n = 29) |
|------------------|---------------------|----------------------|
| Age (yr)         | 46.3 ± 17.8         | 42.0 ± 19.3          |
| Gender (M/F)     | 22/8                | 24/5                 |
| Height (cm)      | 163.2 ± 15.0        | 168.8 ± 8.3          |
| Weight (kg)      | 66.5 ± 15.1         | 64.7 ± 11.2          |
| ASA (I/II/III)   | 14/13/3             | 16/10/3              |
| Mallampati score | 1/2/3/4             | 5/20/5/0             |
| Airway device size | 7/20/3              | 8/14/7/0             |
| LMA (3/4/5)      |                     |                      |
| SLIPA (47/49/51/53/55) | 1/2/11/13/2        |
| Duration of anesthesia (min) | 131.7 ± 76.8 | 131.3 ± 71.9 |

Values are mean ± SD or number. PLMA: laryngeal mask airway ProSeal™, SLIPA: streamlined liner of the pharynx airway.

Fig. 1. The SLIPA in detail. The SLIPA is made of soft hollow plastic. Because the SLIPA resembles the anatomy of the pharynx, an inflatable cuff is unnecessary. Instead, the body serves as a reservoir for regurgitated liquids.
A comparison of SLIPA and PLMA

Vol. 58, No. 5, May 2010

only vocal cords seen, 4 points; vocal cords and back of the
co., Japan), hypopharynx placement was scored as follows:
fiberoptic bronchoscope (Olympus BF-3C40, Olympus optical
valve locked, and fresh gas flow 3 L/min supplied [4]. Using a
when the mechanical ventilation was stopped, the pop-off
the anesthetic circuit by a pressure gauge did not rise anymore
highest peak reached when the airway pressure measured in
was rotated 60 degrees to the right and left to see if there was
To evaluate if the airway was appropriately secured, the head
was performed for 1 minute and a second attempt was made
with a different size. If that also failed, then the patient was
excluded from the study, and the surgery was performed by
endotracheal intubation. In the process of insertion, the
insertion duration (from the point of starting placement to the
point of successful mechanical ventilation), and the number
of insertion attempts were measured. The level of difficulty of
insertion (easy, moderate, difficult) was noted subjectively by
the anesthesiologist who inserted the device.

The same anesthetic machines (Astiva/5, Datex-Ohmeda
Inc., USA) were used throughout the study. Ventilation during
the anesthesia was performed with a tidal volume of 10 ml/kg,
respiratory rate of 12/min and the inspiratory: expiratory ratio
was 1 : 2.

To assess the hemodynamic response to insertion of the
device, systolic blood pressure (SBP), diastolic blood pressure
(DBP), mean blood pressure (MBP), and heart rate (HR) were
recorded 5 minutes before and after the insertion at 1 minute
intervals. The maximal change in mean value (%) was cal-
culated by the ratio of the mean value immediately before
insertion to the highest mean value immediately after insertion.
During insertion, if the blood pressure decreased by more than
20% of the blood pressure measured in the ward, ephedrine 5
mg was administered.

To evaluate if the airway was appropriately secured, the head
was rotated 60 degrees to the right and left to see if there was
any leakage. Also, oral leak pressure (OLP) was measured as the
highest peak reached when the airway pressure measured in
the anesthetic circuit by a pressure gauge did not rise anymore
when the mechanical ventilation was stopped, the pop-off
valve locked, and fresh gas flow 3 L/min supplied [4]. Using a
fiberoptic bronchoscope (Olympus BF-3C40, Olympus optical
c., Japan), hypopharynx placement was scored as follows:
only vocal cords seen, 4 points; vocal cords and back of the
epiglottis seen, 3 points; vocal cords and front of epiglottis seen,
2 points; vocal cords not seen, 1 point [5]. At the beginning and
end of ventilation, gastric distension was checked for with a
stethoscope and the naked eye. 10 minutes after the insertion, a
spirometer module (S/5 Compact anesthesia monitor [6], Datex-
Ohmeda, USA) was used to measure the maximum inspiratory
pressure, plateau airway pressure (Pplat), mean airway pressure
(MAP), peak end expiratory pressure (PEEP), airway resistance
(Raw), and compliance (Cdyn). The spirometer module was
attached to the patient's airway device as a side stream which
allowed for extraction of respiratory gas. After the initial eval-
uation, the mid-surgery tidal volume and respiratory rate were
controlled to achieve an end tidal carbon dioxide value of 35–
40 mmHg.

After the surgery, the airway device was removed and evidence
of blood stains or regurgitation checked for. In the recovery
room, one anesthesiologist, who did not know which group
the patient belonged to, evaluated the postoperative sore
throat pain on the visual analogue scale (VAS score, 0–10) and
checked for other complications.

To calculate the sample size required for this study, time taken
for intubation in a similar clinical setting was taken account [3].
A two-tailed α error of 5% and a β error of 20% was accepted
in the detection of differences of 4 second in the time taken for
intubation between the two groups. Based on this calculation,
the required size per group was 30.

Normally distributed data were compared between groups
by unpaired t-testing, and presented as mean value ± standard
deviation and presents mean value ± standard error in the
figure. Descriptive variables were analyzed, as appropriate,
by Chi-square test or Fischer's exact test. The blood pressures
and heart rates between groups were compared using two way
repeated measures ANOVA. Within a group, comparison with
the base data was performed by one way repeated measures
ANOVA. P values less than 0.05 were considered statistically
significant, and statistical analysis was performed with
Sigmastat (version 2.03, SPSS Ins., USA).

**Results**

The success rate on the first attempt for the 2 groups is shown
in Table 2. When failed on the first attempt, all second attempts
were successful in the PLMA group. Both attempts failed in one
patient in the SLIPA group; this patient was thus excluded from
the study and endotracheal intubation performed. In the SLIPA
group, insertion time was significantly longer (P < 0.05, Table 2).

The hemodynamic response to insertion is shown in Fig. 2.
The maximal change of mean value (%) for the PLMA group
was 1.4% and 7.6% for HR, 1.4%
and 12.8% for SBP, 1.0% and 16.4% for DBP, and 0.9% and 13.4% for MBP. Ephedrine was administered during the first 5 minutes after insertion if blood pressure fell by more than 20% of the preoperative ward blood pressure. This occurred in 3 PLMA cases and 1 SLIPA case.

The mean value of OLP in the PLMA group was 26.1 cmH\textsubscript{2}O. In the SLIPA group, it was 24.5 cmH\textsubscript{2}O. This difference was not significant. There was also no significant difference in gas leakage depending on the head position. After insertion, fiberoptic bronchoscopy findings of the hypopharynx of the two groups were not different. Gastric distension was not observed in either group before or after surgery (Table 2).

Lung mechanics data were similar in the two groups. Both devices supplied adequate ventilation (Table 3).

After the end of the surgery, blood stains were found on the airway device in 12 patients in the SLIPA group (40%), and in 2 people in the PLMA group (6.7%). This difference was statistically significant (P \textless 0.05). Complications observed in the recovery room included sore throat, which was seen in 46.7% (n = 14) in the SLIPA group and 26.7% (n = 8) in the PLMA group. However, this difference was not significant. The VAS scores of

**Table 2. Airway Observations**

|                              | PLMA group (n = 30) | SLIPA group (n = 29) |
|------------------------------|---------------------|----------------------|
| First insertion success rate (%)* | 93.3 (n = 28)       | 73.3 (n = 22)        |
| Insertion time (sec) * †      | 7.3 ± 3.2           | 10.5 ± 3.6           |
| Easy of insertion             | 13/16/1             | 9/9/11               |
| (easy, normal, difficult)*    |                     |                      |
| Oropharyngeal leak pressure (mmHg) | 26.1 ± 8.8         | 24.5 ± 6.7           |
| Leakage (Rt/Lt)               | 7/7                 | 9/7                  |
| Fiberoptic bronchoscopic view | 2/8/15/4            | 4/14/10/2            |
| (1/2/3/4)                     |                     |                      |
| Gastric distension (pre/post) | 0/0                 | 0/0                  |

Values are mean ± SD or number. PLMA: laryngeal mask airway ProSealTM, SLIPA: streamlined liner of the pharynx airway, pre: pre-operation, post: post-operation. *Significantly (P < 0.05) different between groups. † Mann-Whitney U test because of abnormal distribution.

**Fig. 2.** Hemodynamic responses to device insertion in the two groups. (A) Systolic blood pressure (SBP), (B) Diastolic blood pressure (DBP), (C) Mean arterial pressure (MAP), (D) Heart rate (HR). SLIPA: streamlined liner of the pharynx airway, PLMA: laryngeal mask airway ProSealTM, pre: just before insertion. Data are expressed as mean ± SEM, *P < 0.05 compared with baseline value, † P < 0.05 compared with PLMA.
sore throat and the incidence of other complications were not significantly different (Table 4).

**Discussion**

An SGA is relatively easier to insert than an endotracheal tube and they are considered for patients in whom intubation is difficult, or for patients in emergency situations [6]. Markus et al. [7] compared classic LMA and SLIPA and had first insertion success rates of 100% and 98% respectively. Similarly, Miller et al. [8] reported first insertion success rates for PLMA and SLIPA of 96% and 98% respectively. In the present study, the first insertion success rates for PLMA and SLIPA were 93.3% and 73.3% respectively. When the first attempt failed, all second attempts were successful in the PLMA group. Both attempts were unsuccessful in one patient in the SLIPA group. Unlike the PLMA, which achieves a pharyngeal seal through post-insertion cuff inflation, the SLIPA comes in a fixed preformed shape. Correct size selection is therefore important for successful SLIPA insertion. In the present study, most first attempt failures were due to inappropriate size selection. When chosen correctly, the second attempt was successful. Previous studies outside of Korea [2,3,7,8] selected SLIPA size by matching the transverse diameter with the patient’s thyroid cartilage. However, in our pilot tests, the SLIPA size chosen by this method was too large and insertion failed in many cases. We therefore tried the size-rule provided by the SLIPA manufacturer, which involved gender and height. However, this rule was too wide range and selection areas often overlapping. It was thus difficult to select the right size on the first attempt. To improve the first attempt success rate, development of SLIPA selection guidelines for Asians is clearly needed. We plan to develop such guidelines in the future.

In the present study, compared to PLMA insertion, SLIPA insertion took longer and was more difficult. We assume this is due to both our relative lack of experience with SLIPA and the difficulty of selecting an appropriate size. Malik et al. [9] found that inexperienced medical students who used Macintosh laryngoscopes for endotracheal intubation had a slow learning curve. It took them a long time to reach a competent level of maneuvering and this signifies that in times of emergency, endotracheal intubation can fail or can cause complications. In contrast, according to the research of Reinhart et al. [10], the LMA (the most common SGA) has a shorter insertion time, higher success rate on the first attempt, and greater ease of use than endotracheal intubation. In a study on medical students with no experience of the SLIPA, SLIPA success rates and preference rates were higher than those of SS-LMs (soft seal laryngeal masks) and the SLIPA had a shorter insertion time [11]. In addition, because it did not require extra handling like filling the cuff or using the strap, it was more convenient to use. It therefore seems likely that, as with the LMA, the SLIPA can take the place of endotracheal intubation for inexperienced users or in times of an emergency.

An advantage of SGA is that it removes the need for a laryngoscope during insertion, thus avoiding direct irritation of the trachea. During endotracheal intubation, a temporary elevation of blood pressure, tachycardia and arrhythmia may occur. Such cardiovascular changes are caused by irritation to the upper airway by the laryngoscope blade, and to the trachea by the intubation of the endotracheal tube and the expansion of the endotracheal tube cuff [12]. Airway management by PLMA is reported to decrease hemodynamic responses compared with endotracheal intubation [13]. Puri et al. [3] reported that in SLIPA use, blood pressure and heart rate increase more than in PLMA uses, but this difference was not as significant as the difference in the use of endotracheal intubation. In this study, the PLMA group showed steady hemodynamic responses before and after insertion, whereas in the SLIPA group, 1 minute after insertion, the blood pressure and heart rate significantly increased to the maximum value. The longer the insertion time, the greater the risk of tachycardia and hypertension [14]. Good SLIPA insertion technique and correct size selection can thus reduce insertion time and produce a more stable hemodynamic response. The maximal change of mean value (%) in blood pressure and heart rate during endotracheal intubation in

| Table 3. Lung Mechanics Data | PLMA group (n = 30) | SLIPA group (n = 29) |
|-----------------------------|---------------------|---------------------|
| PIP (cmH2O)                | 14.7 ± 4.6          | 13.7 ± 3.1          |
| Pplat (cmH2O)              | 12.4 ± 2.3          | 12.4 ± 1.8          |
| MAP (cmH2O)                | 6.3 ± 1.6           | 6.3 ± 0.9           |
| PEEP (cmH2O)               | 2.9 ± 0.7           | 3.1 ± 0.4           |
| Cdyn (ml/cmH2O/l/s)        | 61.8 ± 17.6         | 63.9 ± 10.6         |
| Raw (cmH2O/l/s)            | 9.0 ± 5.9           | 7.2 ± 4.8           |

Mean ± SD. PLMA: laryngeal mask airway ProSeal™, SLIPA: streamlined liner of the pharynx airway. PLMA: laryngeal mask airway, PIP: peak inspiratory pressure, Pplat: plateau airway pressure, MAP: mean airway pressure, PEEP: peak end expiratory pressure, Cdyn: compliance, Raw: airway resistance.

| Table 4. Complication | PLMA group (n = 30) | SLIPA group (n = 29) |
|-----------------------|---------------------|---------------------|
| Blood stain*          | 2                   | 12                  |
| Regurgitation         | 0                   | 0                   |
| Sore throat           | 8                   | 14                  |
| VAS (sore throat)     | 1.0 ± 2.2           | 1.9 ± 2.7           |
| Nausea/Vomiting       | 2/1                 | 3/1                 |

Values are mean ± SD or number. PLMA: laryngeal mask airway ProSeal™, SLIPA: streamlined liner of the pharynx airway. *Significantly (P < 0.05) different between groups.
the report by Kim et al. [13] was 55.7% for SBP, 47.5% for DBP, and 48.2% for HR. According to Lee et al. [15], these values were 51%, 58%, and 48% respectively. In the present study the respective values for the SLIPA group were 12.8%, 16.4%, and 7.6%. Although not directly measured in this study, SLIPA insertion in comparison to endotracheal intubation is assumed to bring fewer hemodynamic changes.

When positive pressure ventilation using LMA, gas leakage and gastric distension are caused by high airway pressure or the inappropriate location of the laryngeal mask. The LMA maintains the airway though a firm seal between its cuff and the structures surrounding the glottis. In contrast to endotracheal intubation, a PIP higher than the OLP can thus result in gas leakage around the cuff. This can trigger inadequate ventilation or gastric distension, heightening the risk of regurgitation and pulmonary aspiration and increasing the incidence of postoperative nausea [16]. During LMA use, gas leakage and gastric distension have been reported when positive pressure ventilation was used with inspiratory gas pressures of over 20 cm H2O [16]. Devitt et al. [17] reported gastric distension in 35% of patients at 30 cmH2O airway pressure. The PLMA was designed so that the larger, wedge-shaped cuff would plug gaps in the proximal pharynx, and it forms a more effective seal with the upper airway than the conventional LMA [18]. In a comparative study of OLP in PLMA and LMA usage [19], the PLMA had a 10 cmH2O higher OLP than the LMA. In the present study there was no significant difference in OLP between the PLMA and the SLIPA, and both groups had a higher OLP than the mean value of PIP during surgery. Furthermore, gas leakage and gastric distension were not observed during surgery, so in our judgment the risk of pulmonary aspiration caused by both devices was not high. Also, there was no significant difference in gas leakage associated with head position in the two groups. Both groups had relatively safe airway maintenance.

In addition, if the SGA in placed incorrectly it cannot seal effectively with the periglottic tissue and gas leakage may result. The leaked gas will then flow into the esophagus and cause gastric distension and pulmonary aspiration. As assessed by fiberoptic bronchoscopy, a hypopharynx placement score of 3 or 4 points is considered to indicate appropriate insertion location. Gaitini et al. [20] reported 60% of the LMAs as being in the appropriate location. When PLMAs and LMAs were compared, 48% of PLMAs and 74% of LMAs were reportedly successful. This means LMAs were easier to insert [21]. In the present study, 60% of PLMAs and 40% of SLIPAs were deemed to be appropriately inserted at fiberoptic assessment. However, there are reported cases that the fiberoptic position was thought be accurate and effective ventilation was assumed to be achieved, but the actual location was improper [22]. Therefore, fiberoptic findings cannot completely exclude the possibility of gastric distension. However, in the present study gastric distension was not observed before or after surgery. Lung mechanics data were the same for both SLIPA and PLMA, and they both gave adequate ventilation.

After removal of the airway devices, SLIPAs had significantly higher incidence of blood stains than PLMAs. The PLMA is made of a soft material with its position secured by insertion of air into the cuff, whereas the SLIPA is made of stiffer plastic material than the PLMA and is of a fixed shape. The SLIPA thus causes more direct trauma to the oral mucosa. In a comparison of postoperative complications in the recovery room, PLMAs have been reported to have fewer cases of sore throat compared to endotracheal tubes and a lower severity of pain [23]. The incidences of sore throat according to Miller et al. [8] were 57%, 49%, and 30% respectively for endotracheal tubes, SLIPA, and PLMA. The present study yielded similar results, with the SLIPA group having a relatively high incidence of sore throat (though without statistical significance), the VAS scores in more than half of the cases being below 3, and the severity of sore throat being mild. Compared to PLMAs, SLIPAs had more cases of sore throat, attributable to the stiff material the SLIPA is made of. The material can irritate and damage the pharyngeal mucosa, and it also causes a higher number of insertion attempts. Nausea, vomiting, and the other complications were uncommon in both groups.

When a cuff is used to seal the pharynx, as is the case with PLMAs, the initial cuff pressure of 60 cm H2O gradually increases because of N2O diffusion across the cuff wall. The intra cuff pressure significantly increases over 30 minutes [24], increasing the risk of ischemic damage of the laryngopharyngeal mucosa. It is thus important to constantly monitor the intracuff pressure [25]. Additionally, the LMA cuff can also cause pressure neuropraxia of the lingual nerve, hypoglossal nerve, and recurrent laryngeal nerve, while N2O use can cause an increase incompressive and/or stretching forces on oropharynx [26]. In the present study, usage of the PLMA was also associated with elevation of cuff pressure during surgery, with adjustment of the cuff pressure to 60 cmH2O with a hand pressure gauge necessary on several occasions. Furthermore, in 2 cases the change in cuff pressure caused PLMA slippage and hence gas leakage, with the PLMA thus requiring repositioning. The advantage of SLIPAs is that they do not have cuffs, and thus are unaffected by the use of N2O.

The PLMA is designed for multiple sterilizations and reuse. The manufacturer explains that autoclaving can be performed up to 40 times. However, washing and autoclaving cannot completely remove protein deposits. Many different methods for cleaning and sterilizing have thus been discussed, but complete removal of protein is known to be impossible [27]. Protein contamination of reused medical devices can cause...
the transmission of prion proteins, with PLMA reuse running the risk of aiding in the transmission of variant Creutzfeldt-Jacob disease. Also, when many PLMAs are washed together, protein cross-contamination can occur [28]. For such reasons, using disposable devices which have no difference in effectiveness lowers the risk of contamination and is more hygienic. As SLIPA is cheaper than PLMA, disposal after single use is advised. A further economical advantage is the lower cost of cleaning and sterilization.

As a limitation of the present study, while we have much experience and skill at PLMA insertion, we have comparatively little experience with using SLIPA. Puri et al. [3] stated that SLIPA had a short learning curve, with a plateau at 13 cases. In the present study, all the SLIPA insertions were performed by one anesthesiologist who had experience of more than 20 SLIPA insertions before this study. The time taken for SLIPA insertion and the number of attempts required both decreased as the study progressed (Fig. 3). In contrast, the PLMA insertion time and required attempts did not change (Fig. 3). We assume this difference to be due to accumulated experience and accept that it may have affected the results of the study. Another limitation was that evaluation in the recovery room was performed by a blinded anesthesiologist while all other evaluations were performed by the anesthesiologist who performed the insertions. This may have affected the results. A limitation of the present study, while we have much experience and skill at PLMA insertion, we have comparatively little experience with using SLIPA. Puri et al. [3] stated that SLIPA had a short learning curve, with a plateau at 13 cases. In the present study, all the SLIPA insertions were performed by one anesthesiologist who had experience of more than 20 SLIPA insertions before this study. The time taken for SLIPA insertion and the number of attempts required both decreased as the study progressed (Fig. 3). In contrast, the PLMA insertion time and required attempts did not change (Fig. 3). We assume this difference to be due to accumulated experience and accept that it may have affected the results of the study. Another limitation was that evaluation in the recovery room was performed by a blinded anesthesiologist while all other evaluations were performed by the anesthesiologist who performed the insertions. This may have affected the results.

In conclusion, as a newly developed SGA, SLIPA, compared to PLMA, showed statistically significant differences in ease of insertion and low hemodynamic response to insertion. However, the differences are of an acceptable level and there were no differences in airway maintenance and ventilatory efficiency. SLIPA usage was unaffected by N₂O the lower cost of the SLIPA means it is suitable for single-use. We expect SLIPA usage to replace PLMA usage and the SLIPA will be used widely.

**Fig. 3.** Time taken for insertion (seconds) according to case numbers. SLIPA: streamlined liner of the pharynx airway, PLMA: laryngeal mask airway ProSeal™.

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