Comparison of ProSeal™ laryngeal mask airway (PLMA) with cuffed and uncuffed endotracheal tubes in infants

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ABSTRACT

We aimed to compare cuffed and uncuffed endotracheal tubes (ETTs) with ProSeal™ laryngeal mask airway (PLMA) in terms of airway security and extubation, starting out from the hypothesis that PLMA will provide alternative airway safety to the endotracheal tubes, and that airway complications will be less observed. After obtaining approval from the local Ethics Committee and parental informed consent, 120 pediatric patients 1-24 months old, American Society of Anesthesiologists physical status I-II, requiring general anesthesia for elective lower abdominal surgery, were randomized into PLMA (Group P, n = 40), cuffed ETT (Group C, n = 40), and uncuffed ETT (Group UC, n = 40) groups. The number of intubation or PLMA insertion attempts was recorded. Each patient’s epigastrium was auscultated for gastric insufflation, leak volumes and air leak fractions (leak volume/inspiratory volume) were recorded. Post-operative adverse events related to airway management were also followed up during the first post-operative hour. Demographic and surgical data were similar among the groups. There were significantly fewer airway manipulations in the Group P than in the other groups (\(p < 0.01\)), and leak volume and air leak fractions were greater in the Group UC than in the other two groups (\(p < 0.01\)). Laryngospasm was significantly lower in the Group P during extubation and within the first minute of post-extubation than in the other groups (\(p < 0.01\)). Based on this study, PLMA may be a good alternative to cuffed and uncuffed ETTs for airway management of infants due to the ease of manipulation and lower incidence of laryngospasm.

KEY WORDS: Infant; airway; ProSeal™ laryngeal mask airway; endotracheal tubes; post-operative complications

INTRODUCTION

Elective lower abdominal surgery is frequently performed under general anesthesia among pediatric patient population. Developments in anesthesia practice and airway management have increased pediatric anesthesia safety.

Improved airway management efficiency was sought through the use of cuffed or uncuffed tracheal tubes followed by the use of a laryngeal mask airway (LMA). The advantages of each over the others have changed during the last decade due to developments in design, changes in production material, and the availability of various types.

Anesthetists widely use uncuffed tracheal tubes for pediatric patients of 8-10 years old or younger because of the advantages of good seal at the cricoid rings, decreased pressure, and thus a lower risk for mucosal injury [1]. Cuffed tubes have the advantage of creating a leak-free breathing system during positive pressure ventilation with a lower tracheal tube exchange rate and a decreased risk of aspiration compared to uncuffed tubes [2]. However, subglottic stenosis due to cuff pressure-related mucosal hypoperfusion has been attributed to the use of cuffed tubes, making their use controversial [2]. In contrast, LMA has the advantages of a lack of direct contact with the trachea, no requirement for direct laryngoscopy, and a lower incidence of coughing [3].

The ProSeal™ LMA (PLMA) was introduced to clinical practice with additional advantages over tracheal tubes and LMA. The PLMA incorporates an esophageal drainage tube and a bite block. Several studies have demonstrated that higher inspired pressures are possible with PLMA compared to LMA [4-6]. Furthermore, the drainage tube eases stomach aspiration and prevents distension [7]. Although pediatric sizes of PLMAs are in use, related studies on their efficacy are limited [8].

Our primary endpoint was to compare leak volume and fraction between the PLMA and cuffed and uncuffed endotracheal tubes (ETTs) in 1-24 months old pediatric patients undergoing elective lower abdominal surgery. Secondary endpoints were to compare gastric insufflation, airway device insertion, and post-operative complications.
MATERIALS AND METHODS

Patients

This prospective study was approved by the Ethics Committee of Ankara University Faculty of Medicine (03.05.2010, 10-179) and performed in accordance with the ethical standards outlined in the current version of the Declaration of Helsinki. After obtaining informed consent of the parents, 120 children, age 1-24 months, American Society of Anesthesiologists (ASA) physical status I-II, were included in the study. The patients were scheduled for lower abdominal surgery at Ankara University Faculty of Medicine, Pediatric Surgery Unit (hypospadias, retracted testicles, and inguinal hernia), between June 2010 and June 2011. The 120 patients were randomized into PLMA (Group P, n = 40) group and two endotracheal intubation groups: Cuffed ETT (Group C, n = 40) and uncuffed ETT (Group UC, n = 40).

Premature patients and patients with congenital abnormalities, risk of aspiration, upper respiratory tract infection, acute or chronic pulmonary diseases, or risk for difficult intubation were excluded from the study.

Procedures

Following routine monitoring, unconsciousness was induced with 8% sevothlurane in O₂ and venous access was established with a 24-gauge intravenous catheter. Anesthesia was induced with 0.5 mg/kg ketamine, 3.0 mg/kg propofol, and 1 µg/kg remifentanil. No muscle relaxants were used. The sizes of the tubes and LMA were selected according to the Broselow-Luten classification [9].

Inadvertent use of mask ventilation was avoided during the induction of anesthesia before the insertion of the airway device. After inserting the PLMA in the Group P, the correct position was confirmed by the absence of audible sound escaping from the mouth and by adequate chest expansion during ventilation. A manometer (Portex Cufflator, Endotracheal Tube Inflator and Manometer, Portex® Limited, Hythe, Kent, UK) was used to adjust the intracuff pressure to 40 cm H₂O. Then, gastric insufflation was assessed by auscultation of the epigastrium, which was performed by a blinded observer [10]. Then, a nasogastric tube was inserted through the nose of the patient. Furthermore, air leakage to the stomach was controlled by the use of bubbles (foam) at the proximal end of the nasogastric tube. The number of airway device insertion attempts was recorded in all groups.

Mechanical ventilation was adjusted intraoperatively with a fresh gas flow of 4 L/min, 10 mL/kg tidal volume, and 5 cm H₂O positive end-expiratory pressure. N₂O was not used. The respiratory rate was adjusted for an end-tidal CO₂ of 35-45 mm Hg.

The inspiratory and expiratory tidal volumes were recorded during the first 10 inspiration/expiration cycles in all groups, and the difference between them was calculated as leak volume. The ratio of leak volume to inspiratory volume was recorded as the leak fraction.

During the operation, heart rate, mean arterial pressure, pulse oximetry, and airway pressures (peak and plateau pressures) were recorded. At the end of surgery, all airway devices were removed with the patient asleep.

After extubation, laryngospasm, stridor, crup, vomiting, retching, coughing, blood staining on the LMA or tube, blood at aspiration, treatment of stridor, desaturation (SpO₂ ≤90%), and re-intubation were recorded at minute 1, 3, 5, 10, 20, 30, 45, and 60 by a blinded observer.

Statistical analysis

The power analysis sample size (PASS) estimation of the study was performed using PASS software. From preliminary data, we calculated with alpha set at 0.05 that 40 patients per group would give a statistical power of 82% to detect a 25% difference in the leak fraction between the groups. About 50 patients in each group were recruited because of the possibility of dropouts (Figure 1). SPSS 1.5 software (SPSS, Inc., Chicago, IL, USA) was used for the statistical analysis. Medians were compared using the Kruskal-Wallis analysis of variance and Mann-Whitney U test. The Chi-square test was
used for the group percentage comparisons. The Cochran test was used for the duration in the groups as yes/no fractions. A $p < 0.05$ was considered statistically significant.

RESULTS

A total of 120 patients were randomized in this study: 40 in the Group P, 40 in the Group C, and 40 in the Group UC. The gender and age of the three groups were similar. However, the weight of the Group UC was significantly different from that of Group C ($p = 0.01$). The demographic and surgical data are presented in Table 1.

PLMA sizes between 1.5 and 2.0 were used for 22 (55%) and 18 (45%) patients, respectively. Significant differences were not detected for ETT sizes in the C and UC groups. The ETTs were changed to find the appropriate size in 10 patients in the Group UC and in 7 patients in the Group C.

A significantly fewer number of attempts were made to insert the airway device in the Group P than in the groups C and UC ($p < 0.01$ and $p < 0.01$, respectively). No significant difference was recorded between the groups C and UC ($p = 1.00$).

No significant difference in gastric insufflation was observed between the groups ($p = 0.24$).

A significant difference in leak volumes and leak fractions was observed between the Group UC and the other groups ($p < 0.01$) (Figures 2 and 3).

The hemodynamic parameters and airway pressures (peak and plateau) were not significantly different among the groups.

The incidence of laryngospasm at extubation was significantly lower in the Group P than in the other groups, but no difference was observed between the groups C and UC ($p < 0.01$). The incidence of coughing was lower in the Group P than in the other groups ($p = 0.03$). No differences were observed among the groups related to stridor, croup,
DISCUSSION

Our results show that a PLMA can be used safely in 1-24 months old pediatric patients undergoing lower abdominal surgery and that airway management is easier with fewer attempts to secure the airway. In addition, the incidence of laryngospasm and coughing was lower in the Group P compared to the incidence in the other groups. Furthermore, the PLMA and cuffed tubes had fewer leaks than did the uncuffed tubes.

During positive pressure ventilation, supraglottic devices always present a risk of potential aspiration of gastric contents due to gastric insufflation and decreased barrier pressure. Therefore, the use of ETTs is more reliable for security and management of the airway [13]. When using a classical LMA, pressure-controlled ventilation is preferred to volume-controlled ventilation to decrease gastric insufflation [7]. However, this is not critical when using a PLMA, as the drainage tube protects patients from gastric insufflation [7].

Epigastic auscultation is a reliable technique for detecting gastric insufflation. Epigastic auscultation should be repeated to reduce the false positive rate [14]. In the present study, no difference in the incidence of gastric insufflation was observed among the groups.

Tracheal mucosal injury risk can be prevented using an appropriate ETT size and testing for air leaks [15]. The incidence of gastric insufflation and post-operative complications did not differ for the cuffed and uncuffed ETTS after choosing
the appropriate tube size following the air leak tests and by determining the cuff pressures with a manometer.

High cuff pressures while using a classical LMA decrease mucosal perfusion; thus increasing the incidence of post-operative airway morbidity [2]. Ong et al. reported that a pressure of 60 cm H₂O can be applied to a classical LMA [2]. However, in our study, the PLMA cuff pressure decreased to 40 cm H₂O, and the incidence of airway morbidities was similar. Although tracheal mucosal cuff pressure is not clearly defined for pediatric patients, we used a manometer to control the cuff pressure at 20 cm H₂O in the Group C and no significant difference was observed between the groups C and UC related to airway morbidity.

Engelhardt et al. compared classical LMA and cuffed and uncuffed ETTs in 45 pediatric patients and did not find any difference related to post-operative complications [13]. Similarly, in a multicenter study by Weiss et al., cuffed and uncuffed ETTs were compared in 2,246 pediatric patients, and no differences were found with regard to post-operative complications [15]. However, in the present study, the PLMA was advantageous over the ET Ts with a lower incidence of post-operative laryngospasm and coughing. The 40 cm H₂O limitation on the PLMA cuff pressure was considered to contribute to the decreased incidence of post-operative complications.

Although the use of LMA during positive pressure ventilation in adults is not new, its use in pediatric patients is controversial [16]. Engelhardt et al. showed that a LMA is as efficient as cuffed ET Ts and superior to uncuffed ET Ts during low flow pressure-controlled ventilation [13]. In addition, Wheeler et al. demonstrated that a PLMA can be an alternative to tracheal intubation for positive pressure ventilation [8]. In the present study, the PLMA was as efficient as the cuffed ETT when comparing the peak and plateau pressures and leak volumes during the volume-controlled ventilation.

All airway devices were removed with the patient asleep to observed post-operative complications by the time measurement in the recovery phase. Thus, the use of PLMA caused less laryngospasm in the early period after the extubation.

This study had some limitations. First, the tracheal tubes were changed more often than the devices used in the PLMA group based on the leak criteria defined in this study. This may have influenced the incidence of post-extubation adverse events. In addition, the anatomy of a 1 month old infant is different from that of a 24 months old baby. It may have been better to perform a subgroup analysis based on patient age with a larger patient population.

In conclusion, the PLMA may be a suitable method for airway management of 1-24 months old pediatric patients undergoing lower abdominal surgery. Furthermore, the PLMA was a good alternative to the cuffed and uncuffed ET Ts with easier insertion, adequate positive pressure ventilation, and lower incidence of post-operative laryngospasm.

DECLARATION OF INTERESTS

The authors declare no conflict of interests.

REFERENCES


Eyyup Sabri Ozden, et al.: Comparison of ProSeal™ with cuffed and uncuffed endotracheal tubes
