“Comparison of Cuff Pressure Changes Between Air with Oxygen or Nitrous Oxide with Oxygen During General Anaesthesia Using Proseal LMA for Laproscopic Surgery”

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Abstract

Background:
LMA exerts pressure on the pharyngeal mucosa which may lead to throat discomfort. Nitrous oxide is known to diffuse into air containing cavity. Nitrous oxide use causes increase in proseal LMA(PLMA) cuff pressure was proved, but whether the resulting increased cuff pressure leading to laryngopharyngeal morbidity which is clinically important remains unclear. We therefore, tested the hypothesis.

Methods: Eighty patients are randomly divided into group A(O2+Air) and group N(O2+N2O) each containing 40 Patients using computer generated randomization list. Patients monitored during surgical procedure regarding intraoperative hemodynamic changes, increase in PLMA cuff pressure, number of deflations required and laryngopharyngeal morbidity during intra and postoperatively upto 24 hrs. PLMA cuff pressure was monitored using cuff pressure monitor[VBM Aneroid meter].

Results: There were no significant intraoperative differences between two groups air and nitrous oxide with respect to hemodynamic parameters, but statistically significant (p value < 0.001) cuff pressure changes with nitrous oxide use which was exceeding > 60 cms of H2O and upto six deflations to maintain pressure in PLMA were required. There was no statistically significant difference for laryngopharyngeal morbidity, probably because we have limited cuff pressure upto 60 cms of H2O in nitrous oxide group as well in laproscopic surgeries > 3 hrs.

Conclusions: Our study concludes that use of cuff pressure monitoring in PLMA to maintain cuff pressure as recommended by the manufacturer probably reduces the incidence of postoperative pharyngo-laryngeal morbidity. Cuff pressures are increased with nitrous oxide use and repeated deflation of cuff is required. But pharyngo-laryngeal morbidity can be limited by deflation and monitoring of cuff pressure for nitrous oxide. Nitrous oxide can be safely used for laproscopic surgeries with PLMA for surgeries lasting less than three hours.

Keywords: PLMA, nitrous oxide, cuff pressure, laryngopharyngeal morbidity, laproscopic surgery.

I. Introduction

PLMA has cuff volume and cuff pressure. This PLMA cuff pressure needs to be monitored for better seal of PLMA. Optimum cuff pressure of PLMA prevents dislodgement and aspiration through PLMA. Factors which affect cuff pressure of PLMA are number of attempts for insertion, duration of surgery, weather ventilated spontaneously or IPPV was used, material of PLMA, gases used for maintenance of anesthesia. Similarly these are the factors which may have effect on cuff pressure and laryngopharyngeal morbidity. Laporoscopic surgery leads to physiological changes due to use of Carbon dioxide which also leads to cuff pressure changes in PLMA, PCO2 levels and incidence of postoperative nausea and vomiting. Keeping all these factors in mind we have planned this study, knowing that use of O2+N2O and O2+Air leads to cuff pressure changes in LMA and if this pressure is not monitored, it leads to increased laryngopharyngeal morbidity due to increased pressure on laryngopharyngeal mucosa and decreased mucosal perfusion pressure and possibility of dislodgment of PLMA and aspiration.

II. Methods

After ethics committee approval and informed consent, eighty patients were enrolled in this study.

Inclusion Criteria :
1. Patients undergoing laproscopic surgery.
2. patient of either sex male/female.
3. Age 18-65 yrs.
4. Weight 32 to 82 kg.
5. ASA grading I and II.
Exclusion criteria
1. Inadequate mouth opening
2. Patient with BMI > 35 kg/m2
3. Anticipated difficult airway.
4. Patient having disease with risk of aspiration like GE reflux, hiatus hernia.
5. Oropharyngeal pathology
6. ASA grading III, IV
7. Cervical spine pathology &
8. Pregnancy

Eighty patients were randomly divided into group A and group N containing 40 patients each using computer generated randomization list .

In group A : patients undergo laparoscopic surgery using PLMA with Oxygen+ air.

In group N : patients undergo laparoscopic surgery using PLMA with Oxygen+ nitrous oxide.

After preoperative anesthesia evaluation patients were taken in OR . In OR after securing IV line, standard monitoring was attached. All the patients received Midazolam 0.03 mg/kg, fentanyl 2 µg/kg as premedication. Anaesthesia was induced using propofol 2 mg/kg and vecuronium 0.1 mg/kg. Anaesthesia was maintained using oxygen and air (FIO2 50%), Isoflurane and vecuronium in group A and oxygen and nitrous oxide (FIO2 50%), Isoflurane and vecuronium in group N. A PLMA was (sizes 3, 4 in females and size 4, 5 in males) used as per standard recommendations. The cuff was inflated to a volume as per manufacturer recommendation and pressure was measured which was found to be above 45 cms of water which will be maintained with a maximum upto 60 cms of water throughout the procedure with cuff pressure monitor (VBM aneroid meter). Placement of PLMA was confirmed by Manual ventilation. Expired tidal volume of >8ml/kg, Square wave capnography, no audible leak from the drain tube with peak airway pressures less than 20 cms of water. The gel displacement test done by placing the blob of gel at the tip of drain tube and noting the airway pressure at which it will be ejected. Positive pressure ventilation was started with a tidal volume of 8ml/kg. The time interval between picking up the PLMA and obtaining an effective airway was recorded. In the event of airway obstruction or a significant leak, PLMA was removed and reinserted. A gastric tube (size 14-16) was then passed through the drain tube, case of placement of gastric tube is recorded and its correct placement confirmed by injection of air and epigastric auscultation.

After insertion of suitable proseal LMA (PLMA), cuff was inflated up to recommended volume for that particular LMA. Usually the cuff pressure more than 45 cms of H2O was observed. Hence cuff pressure of 45 cms of H2O was taken as baseline for all the patients. With this pressure adequate seal was maintained and no leak was confirmed. Intraoperatively if cuff pressure was more than 60 cms of water cuff was deflated to 45 cms of H2O.

Group A : Oxygen and air was used for maintenance of anesthesia.

But in this group cuff pressure never crossed 60 cms of H2O throughout the surgery. So no deflation was required in this group.

Group N : Oxygen and nitrous oxide was used for maintenance of anesthesia.

In this group intracuff pressure was crossing limit of 60 cms of H2O. So cuff pressure was deflated to 45 cms of H2O whenever it crossed 60 cms of H2O. Number of deflations and time for deflation were noted. Protocol to maintain SpO2 above 95% and EtCO2 between 35-45 mm Hg was observed by adjusting respiratory rate (RR) and tidal volume. If SpO2 falls below 97% and if EtCO2 did not improve, tidal volume was increased to 10 ml/kg. If EtCO2 increased above 45 mm Hg, RR increased to 14 to 18 breaths/min. Peak airway pressure was recorded once abdominal insufflation started and reached 12 mm Hg and pressures was kept between 12-14 mm Hg. Episodes of gastric insufflation, regurgitation and aspiration were recorded. Intraoperative analgesia was achieved with IV fentanyl 1 µg/kg and IV diclofenac 1 mg/kg for postoperative analgesia. Neuromuscular blockade reversed with glycopyrrolate 8 µg/kg and neostigmine 0.05 mg/kg.

At the end of the surgery, the anesthesiologist removed the PLMA when the patient was awake and opened his/her mouth. Presence of blood on the PLMA was noted. Oral cavity was examined for any injury to lip, teeth, gums, pillars and soft palate. The patients were monitored in the Recovery Room and enquired for laryngopharyngeal morbidity like sore throat, dysphagia, dysphonia and nausea and vomiting. Patients were discharged to the ward when recovery was deemed adequate. The patients were enquired for laryngopharyngeal morbidity after 24 hrs as well.

III. Results

There were no significant intraoperative differences between the two groups A and N with respect to hemodynamic parameters, operating conditions or bowel distension but there were statistically significant (p value < 0.001) cuff pressure changes with nitrous oxide use which was exceeding > 60 cms of H2O (figure 1, 2).
Table 1). If duration of surgery was up to 3 hrs, number of deflation of cuff pressure were up to 5 or more in nitrous oxide group (figure 2). There was no statistically significant difference between the two groups in terms of laryngopharyngeal morbidity, probably because we have limited cuff pressure up to 60 cms of H2O in nitrous oxide group (Table 3, 4, 5, and figure 3, 4, 5, 6).

**Figure 1:** Cuff pressure at various durations in the study groups

**Table no 1:** Comparison among study group for intracuff pressure
Comparison of Cuff Pressure Changes Between Air With Oxygen Or Nitrous Oxide With

Figure 2: Number of deflations for cuff pressure in the study group

Figure 3: Incidence of Immediate postop sore throat in the study groups
Comparison of Cuff Pressure Changes Between Air With Oxygen Or Nitrous Oxide With

Figure 4: Incidence of Immediate postop Nausea in the study groups

Figure 5: Incidence of Immediate postop vomiting in the study groups
Comparison of Cuff Pressure Changes Between Air With Oxygen Or Nitrous Oxide With

Figure no 6: postop upto 24hrs various complications in the study groups

Table 2 : Number Of Cuff Deflations In Study Group And Time.

<table>
<thead>
<tr>
<th>Time in mins</th>
<th>Group A N = 40 Pressure cms H2O</th>
<th>Group N N = 40 Pressure cms H2O</th>
<th>Number of Deflations in groups A N</th>
<th>Total number of deflations at particular time approx in group A N</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>45</td>
<td>45</td>
<td>0 0</td>
<td>0 0</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>46.78</td>
<td>53.75</td>
<td>0 0</td>
<td>0 0</td>
<td>0.000</td>
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<tr>
<td>15</td>
<td>48.58</td>
<td>60.93</td>
<td>0 1</td>
<td>0 1</td>
<td>0.000</td>
</tr>
<tr>
<td>30</td>
<td>49.40</td>
<td>59.40</td>
<td>0 1</td>
<td>0 2</td>
<td>0.000</td>
</tr>
<tr>
<td>60</td>
<td>49.12</td>
<td>60.31</td>
<td>0 2</td>
<td>0 4</td>
<td>0.000</td>
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<td>90</td>
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<td>56.40</td>
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<td>120</td>
<td>45.63</td>
<td>54.35</td>
<td>0 1</td>
<td>0 6</td>
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Table 3 : Laryngopharyngeal Morbidity With Plma

<table>
<thead>
<tr>
<th>Laryngopharyngeal Morbidity With Plma</th>
<th>Group A</th>
<th>Group N</th>
</tr>
</thead>
<tbody>
<tr>
<td>INTRAOPERATIVE</td>
<td>[ 40 ]</td>
<td>[ 40 ]</td>
</tr>
<tr>
<td>1. Leak</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2. Gastric insufflations</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>3. Regurgitation and aspiration</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>AT REMOVAL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Coughing</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>2. Blood staining of device</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>3. Trauma to lip, teeth, tongue</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>POSTOPERATIVE</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Nausea</td>
<td>3</td>
<td>19</td>
</tr>
<tr>
<td>2. Vomiting</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>3. Sore throat</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>4. Dysphagia</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>5. Dysphonia</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>6. Dysarthria</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
Airway devices have cuffs which are permeable to a variety of gases depending on their partial pressure and solubility. The composition and thickness of the cuff material (latex, silicone or polyvinyl chloride) play a significant role in the intracuff pressure changes during anesthesia. Nitrous oxide and other gases diffuse into air filled cuffs of tracheal tubes and supraglottic devices, increasing their volume and pressure. We chose this study because increased intra-abdominal pressure from pneumoperitoneum requires higher airway pressures for adequate pulmonary ventilation, for which the PLMA has proved to be adequate in previous studies. Mean time taken for successful placement of PLMA was 12 secs and 13 secs for groups A and group N respectively. Studies by Cook, Shroff and co-workers (median effective time 15 s) corroborated with our study findings. Sharma and coworkers reported a mean insertion time of 13.51 s and 12 s, respectively. In our study there was no difference in both the groups A and group N in terms of pulse rate, mean arterial blood pressure, SpO2, EtCO2 and peak airway pressures. Both groups maintained adequate oxygenation and ventilation perioperatively. Maltby et al and Sharma et al found no statistically significant differences in SpO2 or EtCO2 between the two groups A and N before or during peritoneal insufflations. The observed oropharyngeal seal pressure for PLMA group was 45 cm of H2O (median), with no clinically audible leak throughout the surgery. The Peak Airway Pressure did not increase beyond the oropharyngeal seal pressure throughout surgery.

Table 4: Incidence of immediate postop dysphagia in the study groups

<table>
<thead>
<tr>
<th>DYSPHAGIA</th>
<th>St Group</th>
<th>Group A</th>
<th>Group N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>Count</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Percent</td>
<td>0.0%</td>
<td>2.5%</td>
</tr>
<tr>
<td>No</td>
<td>Count</td>
<td>40</td>
<td>39</td>
</tr>
<tr>
<td></td>
<td>Percent</td>
<td>100.0%</td>
<td>97.5%</td>
</tr>
<tr>
<td>Total</td>
<td>Count</td>
<td>40</td>
<td>40</td>
</tr>
</tbody>
</table>

Chi-Square test Value df P value
Pearson Chi-Square 1.013 1 0.314
Fisher’s Exact Test 1.000

Table 5: Immediate postop dysarthria in the study groups

<table>
<thead>
<tr>
<th>DYSPHONIA</th>
<th>St Group</th>
<th>Group A</th>
<th>Group N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>Count</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Percent</td>
<td>0.0%</td>
<td>2.5%</td>
</tr>
<tr>
<td>No</td>
<td>Count</td>
<td>38</td>
<td>39</td>
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<tr>
<td></td>
<td>Percent</td>
<td>95.0%</td>
<td>97.5%</td>
</tr>
<tr>
<td>Total</td>
<td>Count</td>
<td>40</td>
<td>40</td>
</tr>
</tbody>
</table>

Chi-Square test Value df P value
Pearson Chi-Square 0.346 1 0.556
Fisher’s Exact Test 1.000

IV. Discussion

Airway devices have cuffs which are permeable to a variety of gases depending on their partial pressure and solubility. The composition and thickness of the cuff material (latex, silicone or polyvinyl chloride) play a significant role in the intracuff pressure changes during anesthesia. Nitrous oxide and other gases diffuse into air filled cuffs of tracheal tubes and supraglottic devices, increasing their volume and pressure. We chose this study because increased intra-abdominal pressure from pneumoperitoneum requires higher airway pressures for adequate pulmonary ventilation, for which the PLMA has proved to be adequate in previous studies. Mean time taken for successful placement of PLMA was 12 secs and 13 secs for groups A and group N respectively. Studies by Cook, Shroff and co-workers (median effective time 15 s) corroborated with our study findings. Sharma and coworkers reported a mean insertion time of 13.51 s and 12 s, respectively. In our study there was no difference in both the groups A and group N in terms of pulse rate, mean arterial blood pressure, SpO2, EtCO2 and peak airway pressures. Both groups maintained adequate oxygenation and ventilation perioperatively. Maltby et al and Sharma et al found no statistically significant differences in SpO2 or EtCO2 between the two groups A and N before or during peritoneal insufflations. The observed oropharyngeal seal pressure for PLMA group was 45 cm of H2O (median), with no clinically audible leak throughout the surgery. The Peak Airway Pressure did not increase beyond the oropharyngeal seal pressure throughout surgery. This is in accordance with the findings of previous studies. There was no incidence of regurgitation or aspiration in either groups in our study. Similar results have been reported by others.

In our study we have observed a significant and progressive increase in intracuff pressure of the PLMA over time when nitrous oxide was used as a part of balanced anesthesia technique for laparoscopic surgery. However, the cuff pressure did not change much when air was used instead of nitrous oxide (P < 0.001). In our study general anaesthesia lasting for more than 3hrs, with gas mixture 50% O2+ N2O, it was observed that the intracuff pressure of PLMA increased from baseline of 45 cm of H2O to more than 60 cms of H2O and needed 1 to 6 times of deflation to baseline pressure depending on the duration of surgery. A similar increase in intracuff pressure from 61 cms of H2O to 123 cms of H2O within two hours of surgery has been reported using the laryngeal tube with a silicone cuff. In our study too, the intracuff pressure remained around 50-55 cms of H2O in group A, while in group N the percentage rise in cuff pressure every 15 - 20 minutes from the baseline was significantly higher (P < 0.001), reaching >60 cms of H2O. The maximum increase at first 10 min was due to the increased pressure gradient at initial low intracuff volume. With the passage of time, it declined as the pressure gradient decreased with further diffusion of nitrous oxide into the PLMA cuff. The rise in the intracuff pressure of the tracheal tubes and supraglottic devices is known to increase the ischemic damage to the surrounding pharyngolaryngeal mucosa. Unlike the tracheal tube cuff which expands within the rigid confines of the tracheal rings, the PLMA cuff inflates in the compliant potential space of the pharynx allowing the cuff walls to match the contours of pharyngeal and laryngeal surfaces. A progressive reduction in the pharyngeal mucosal perfusion has been reported when mucosal

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pressure increases from 25 to 60 mmHg while using a cuffed oropharyngeal airway\(^{(20)}\). The cuffs of CLMA and PLMA exert pressure on the pharyngeal mucosa causing a concomitant decrease of pharyngeal perfusion and increase in the incidence of post-operative complications including sore throat, dysphonia, and nerve damage\(^{(20,18)}\). Study with the PLMA suggested that directly measured mucosal perfusion pressure rarely exceeded 25 mmHg\(^{(19)}\) and therefore, did not increase pharyngeal mucosal injury, while others recommend reducing the cuff volume until it just seals the leak\(^{(21,22)}\). The cuffs of PLMA inflated with maximum recommended cuff volumes, exerted lower pressures predominantly below 15 mmHg on the pharyngeal and hypopharyngeal mucosa. It was also reported that PLMA along with nasogastric tube, induce significantly higher pharyngeal pressures in posterior location when compared to other devices. This may be because of the additional cuff on the posterior part of the PLMA\(^{(12)}\). Higher cuff pressure and higher incidence of postoperative sore throat has also been reported after use of PLMA in children while breathing 50% nitrous oxide and oxygen mixture in comparison to patients whose breathing gases were composed of oxygen and air\(^{(24,25)}\). Carbon dioxide used during the laparoscopic procedures may diffuse into the cuff to increase intracuff pressure. However, it does not contribute to rise in intracuff pressure as the cuff pressure remained unchanged in group A. The rise in intracuff pressure in group N can be attributed to the diffusion of nitrous oxide which is more diffusible than carbon dioxide\(^{(26)}\). The reported incidence of sore throat in group A is 10% and for group N is 5%\(^{(27,29)}\). In our study, the incidence of sore throat was low and comparable between the two groups because we have deflated the PLMA cuff pressure in group N, once it is crossing limit of 60 cm of H2O. In our study maximum duration of surgery was two and half hours duration, we have noted that in Group N the cuff pressure started increasing immediately after pneumoperitonium and as soon as it increased above 60cm of H2O we have deflated it. So the number of deflations were more in group N and there were no much cuff pressure changes in group A as it is increased only up to 55 cm of H2O. Number of cuff deflations required in group N is statistically significant with p value of <0.001. Number of deflation were more in cases where surgery lasted for more than 2 hrs in group N(1-5times). Similarly the incidence of dysphagia and dysphonia was comparable probably because of pressure limitation in group N. As the device is being increasingly used for procedures longer than two hours, vigilance is required during its use and excessive gas should be regularly removed from the cuff.

Limitations of our study- We did not monitor the pharyngeal mucosal pressure or intracuffgas mixture due to the non-availability of the appropriate equipment (microchip sensor or gas analyzer). In our study maximum number of attempts of PLMA insertion were up to 3, so there was no significant relationship with the incidence of postoperative sore throat and number of attempts in our study and exclusively relate sore throat with rise in intra-cuff pressures.

Use of cuff pressure monitor is recommended for initial cuff inflation as well as for intraoperative monitoring during laparoscopic surgery using a nitrous oxide based anesthesia techniques. According to Eduardo Figueredo and Miguel Vivar-Diago\(^{(28)}\) the use of IPPV was the cause for post operative pharyngolaryngeal adverse effects and not the cuff pressure or spontaneous ventilation. Since it has been suggested that the use of nitrous oxide may contribute to bowel distention, evaluated the effects of N2O on operating conditions during laparoscopic cholecystectomy in 50 healthy patients using a double-blind protocol design. For maintenance of anesthesia, patients were randomly assigned to one of two treatment groups: 1 (n = 26) received isoflurane with 70% N2O in oxygen, whereas group 2 (n = 24) received isoflurane in an air/O2 mixture. The surgeon (blinded to the anesthetic technique) estimated the degree of technical difficulty before beginning the operation using a five-point scale. At 15-min intervals throughout the operation, the surgeon was asked to evaluate both “overall operating conditions” and degree of “bowel distension” using independent five-point scales. There were no significant intraoperative differences between the two groups with respect to operating conditions or bowel distension. Thus, N2O had no clinically apparent deleterious effects during laparoscopic cholecystectomy. Akca O Et al \(^{(14)}\) studied Patients scheduled for colon resection were anesthetized with isoflurane and 35% oxygen and 65% nitrous oxide (n = 175) or air (n = 169), results suggest that avoiding nitrous oxide administration during prolonged bowel operations will minimize bowel distension. In our study most of the surgery duration was within 3 hrs and there was no noticeable bowel distension during surgery as verbally asked to the surgeons. Surgeon was blinded to balanced anesthesia technique weather O2+ N2O or O2 + Air was utilised.

V. Conclusion

Our study concludes that use of cuff pressure monitoring in PLMA to maintain cuff pressure as recommended by the manufacturer probably reduces the incidence of postoperative pharyngo-laryngeal morbidity like sore throat, dysphagia and dysphonia. Cuff pressures are increased with nitrous oxide use and repeated deflation of cuff is required to maintain target pressure in cuff of PLMA. Nitrous oxide can be safely used for laparoscopic surgeries with PLMA for surgeries lasting less than 3 hours. Pharyngo-laryngeal morbidity can be reduced by deflation and monitoring of cuff pressure for nitrous oxide use.
of room air and N2O+O2 used for ProSeal LMA cuff inflation on 3:

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