Factors Affecting Postoperative Agitation in Adults Following Functional Endoscopic Sinus Surgery: A Randomized, Double-Blinded Controlled Trial

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Abstract

Background: We have noticed a high incidence of postoperative agitation (POA) following functional endoscopic sinus surgery (FESS), we reviewed our anesthesia protocol and designed this study. The present study investigated the possible contributing factors in the development of POA in adults, by avoiding these factors postoperative agitation can be minimized.

Methods: 140 ASA 1 and 2 patients were randomly allocated to either premedication group that received Phenergan 25 mg IM and atropine 0.5mg IM or none premedication group that received an equal volume of water for injection intramuscular injection. Hypotensive anesthesia was induced by nitroglycerin 5 to 20 mcg/kg/minute. In postoperative care unit (PACU), patients were assessed for both agitation and pain using Richmond agitation sedation scale and numerical rating scale respectively. Data were analyzed and compared between groups, P<0.05 was considered significant.

Results: Agitation was more frequent in the premedication group; premedicated patients showed a higher incidence of agitation versus the control group 82% versus 35% respectively P<0.0001. Postoperative agitation was more frequent in young ages, males, smokers and was correlated with pain. No correlation was found with BIS value or postoperative O2.

Conclusion: premedication with atropine and Phenergan, Pain, smoking, young age, male gender was identified as risk factors for the development of postoperative agitation in adults following FESS.

Keywords: Agitation postoperative; Premedication atropine; Premedication Phenergan; Endoscopic sinus surgery
Introduction

In our previous report on postoperative agitation in adults, we have investigated the role of magnesium sulfate as a potential treatment for postoperative agitations following functional endoscopic sinus surgery [1]. However, the high incidence of postoperative agitation in our protocol begs the search for the possible contributing factors to this phenomenon. Postoperative agitation although short-lived it could be harmful to both patients and recovery staff [2,3]. An agitated patient has the potential for self-injury by removing intravenous cannulas, tubes, oxygen masks, and nasal packs. Furthermore, very agitated patients can pose an immediate danger to operating room staff [4]. In the last few years, and during our work with patients subjected to endoscopic sinus surgery, we have observed a high incidence of postoperative agitation that was not terminated by administration of narcotic analgesics. In the postoperative care unit, we noticed some agitated patients forcibly removing their venous and arterial cannulas, Oxygen masks resulting in bleeding and hypoxia. We also recorded bruises and ecchymosis from the restraints of agitated patients. These complications have forced us to review our anesthesia protocol and implement a study to identify the risk factors that might account for this phenomenon. Kim, et al. [5] reported that postoperative agitation following nasal surgery could occur due to several factors; young age, smoking, sevoflurane anesthesia, postoperative pain, the presence of a tracheal tube or a urinary catheter. Some of the drugs that are used in anesthetic pre-medication have been shown to increase excitation and agitations [6]. Amongst these anticholinergic and antihistaminic drugs are incriminated [7]. We have sought this study to explore whether the used Atropine and Phenergan have contributed to the high incidence of postoperative agitation. The other aim is to discover the other contributing factors that are responsible for postoperative agitation in adults following functional endoscopic sinus surgery.

Methods

Patients and Design

This study was approved by the Menofia University ethics committee on January 6th, 2014. 140 American society of anesthesiologist- physical status I and II patients of both genders were enrolled in the study, then written consent was obtained from each participant. Inclusion criteria were age between 18 and 60 years of both sexes, ASA I and II statuses, and patient acceptance to participate in the study. Exclusion criteria were hypertensive patients, cardiac ischemia, cerebrovascular insufficiency, neuromuscular diseases, pregnancy, prolonged treatment with calcium channel blockers, diabetic neuropathy. Patients were randomized by a research randomizer computer system and were assigned to either Pre-medication (Group A) or the control (Group B) by central randomization through phone calls to an anesthesia technician who was not participating in the study; he was furnished patient ID, body weight, and group allocation and was tasked to prepare colorless coded solutions contained in transparent syringes. Group A patients were prem edicated by atropine sulfate Boden Centra limited (0.5 mg/ml) 0.5mg IM and Phenergan (Promethazine HCl, Lab Renaudine, France) as a sedative and antiemetic 25 mg IM one hour before entering the operation theater. Group B patients were given an equal volume of plain water injection intramuscularly. All patients underwent functional endoscopic sinus surgery at Menofia University Hospitals ENT room. For the purpose of estimating the incidence of agitation in relation to age, all patients were classified into 2 categories depending on their ages. Category one included age range from 18 to 30 (Cat 1), category two included age range from 31 and up (Cat 2). The smoking history was taken; smokers were ranked according to the duration and daily consumption of cigarettes into 4 categories 0, 1, 2 and 3. A smoking index was implemented by multiplying the number of cigarettes per day by the smoking years, and the resulted number was graded on a four-point scale: nonsmoker, light, heavy and massive smoker as follows: 0=0, 1= 1:200, 2=201:400 and 3=401:600. Below the smoking index variable, each patient was assigned a number depending on his smoking status, 0= nonsmoker, 1= light smoker, 2= heavy smoker, and 3= massive smoker.

Anesthesia Technique

A large bore 18-gauge intravenous cannula was inserted, all patients underwent general endotracheal anesthesia and were anesthetized by Propofol 2.5 mg/kg, Fentanyl 1 mcg/kg, Atracurium 0.5mg/kg was given to facilitate intubation and ventilation. Anesthesia was maintained by sevoflurane 2% in 50% air and Oxygen. All patients were monitored by standard monitors including, the pulse oximeter, non-invasive blood pressure, end-tidal CO₂, and electrocardiogram. Moreover, an arterial line was inserted, and invasive blood pressure was monitored.
throughout the procedure. Sevoflurane, O₂ percentage concentrations were monitored throughout the procedure. Anesthetic depth was monitored using bispectral index (Aspect XP, USA) to exclude awareness as a contributing factor to the development of agitation. Hypotension was induced by infusion of nitroglycerine (5mg/ml Hospira UK Limited) at a rate of 5 to 20 mcg/ kg /minute titrated to target mean blood pressure of 55±5. The purpose of hypotensive anesthesia was to reduce bleeding and improve the surgical field. After the onset of hypotension Mean arterial pressure and Bispectral index were recorded simultaneously every 10 minutes. The medians of the obtained readings were calculated for each patient and represented the Med. MAP and Med. BIS respectively. Hypotensive anesthesia was maintained throughout the surgical procedure. After the conclusion of surgery, patients were given ephedrine hydrochloride (Martin Dale Pharmaceuticals) 10 mg IV to restore the mean arterial pressure back to the preoperative value. Upon attaining spontaneous breathing patients were given reversal to muscle relaxants and extubated in a wake mode. Upon recovery, one gram Paracetamol (Perfalgan, Bristol Mayers Squeeb) was given intravenously to each patient.

Patient Assessment

Patients were transferred to PACU for observation and assessment for agitation and pain. Upon arrival to PACU, a pulse oximeter and NIBP were attached to the patient. Discharge criteria from PACU were stable vital signs, pain score less than or equal to 2, no nausea or vomiting, calm and alert patient. Patient age, body weight, gender, and vital signs were recorded and compared between groups.

Assessment of agitation

Agitation was defined as a purposeless excessive motor activity and/ or inconclusive sounds or aggressive behavior of the patient with no response to commands. Postoperative agitation was assessed in the PACU by a blinded observer using the Richmond Agitation Scale (RASS) as follows:
0= Alert and calm
+1= Restless; Anxious and /or apprehensive but movements not aggressive.
+2= agitated; Frequent non-purposeful movement.
+3= very agitated; Pulls on or removes the tubes or catheters or has aggressive behavior toward staff.
+4= combative; overly combative or violent.
Total agitation was defined as the algebraic sum of agitation values observed at the measured time points, and this was used to correlate agitation with other variables. The range of agitations was 0= no agitation to 20= severe persistent agitation.

Assessment of pain

Pain scores were assessed using numerical rating scale where 0= no pain and 10 is the worst imaginable pain as rated by the patient. A blinded team member assessed pain after evaluation of last agitation point (30 minutes of recovery). Patient that exhibited no agitation were also evaluated for pain after 30 minutes of recovery. Pethidine was used as rescue analgesic to ameliorate pain. The use of pethidine was intended after evaluation of agitation and assessment of pain to avoid confounders to the outcome measurement. Pethidine was given in 25 mg increments every 10 minutes until pain score ≤ 2.

Statistical analysis

Power analysis was done based on our pilot study comparing the frequency of agitation 10 minutes postoperatively as a primary determinant of sample size. Using means difference between the two groups (effect size) of 25%, given α probability =0.05, power 95% revealed a total sample size of 132 is required, 66 per group to detect 25‰ reduction in postoperative agitation. We increased the total sample size to 140 to avoid a drop-in sample size due to possible exclusions. Descriptive statistics were used to represent patient characteristics. Although the Richmond Agitation Sedation Scale values are ordinal variables, we have used 5 defined points that can be treated as a continuous variable [8]. Continuous variables were tested for normality and were represented by means or medians as appropriate; categorical data were represented by percent of the total. All agitation points were compared by Man-Whitney U test, Medians of pain scores were compared by Mood's test. Correlation between agitation and other variables was evaluated by Spearman correlation test and Chi-square test. The coefficients of determination (R²) for pain, BIS, smoking index and PACU stay were estimated by ANCOVA. P<0.05 was considered significant.

Results

Figure 1 shows, patient enrollment, and randomization process. 140 patients were randomly allocated to premedication group (N=70) or the control group (N=70). 3 patients were excluded from the final analysis because of atypical hypotension 2 in pre-
medication and 1 in the control group. 137 patients were analyzed, 68 in pre-medication and 69 in the control group. There was no difference in patient characteristics (Table1).

<table>
<thead>
<tr>
<th>Variable</th>
<th>Control G</th>
<th>Premedication G</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>68</td>
<td>69</td>
<td></td>
</tr>
<tr>
<td>Age(years)</td>
<td>35±10</td>
<td>33±12</td>
<td>0.1</td>
</tr>
<tr>
<td>BMI (Kg/m2)</td>
<td>30.6±3</td>
<td>31±4</td>
<td>0.06</td>
</tr>
<tr>
<td>ETCO₂ (mm Hg)</td>
<td>36±3</td>
<td>35±4</td>
<td>0.05</td>
</tr>
<tr>
<td>BIS value</td>
<td>40±3</td>
<td>39±2</td>
<td>0.05</td>
</tr>
<tr>
<td>Recovery SPO₂</td>
<td>99±0.9</td>
<td>99±0.8</td>
<td>0.9</td>
</tr>
<tr>
<td>Pre-MAP (mm Hg)</td>
<td>89±4</td>
<td>88.6±4</td>
<td>0.9</td>
</tr>
<tr>
<td>Med MAP (mmHg)</td>
<td>53±3</td>
<td>54±4</td>
<td>0.5</td>
</tr>
<tr>
<td>Rec. MAP (mmHg)</td>
<td>89±4</td>
<td>88.6±4</td>
<td>0.9</td>
</tr>
<tr>
<td>Gender M/F</td>
<td>29/40</td>
<td>30/38</td>
<td></td>
</tr>
</tbody>
</table>

Table 1: Patient characteristics.

Table 1 reveals the patient characteristics, bispectral index, end-tidal carbon dioxide, recovery oxygen saturation and mean arterial blood pressure in both groups. There was no statistical difference between the two groups in these parameters. BMI= body mass index, G= group, ETCO₂=end-tidal carbon dioxide, Recovery SPO₂= Oxygen saturation measured at recovery room, M/F= Male and female.
The incidence of agitation among all patients was found to be 68%; Premedicated patients showed a higher incidence of agitation versus the control group 82% versus 35% respectively P<0.0001. Young patients showed higher incidence and more severe agitation than older patients did. Cat1 patients (n=68) showed an 85% incidence of postoperative agitation (POA), Cat2 (n=69) showed an 35% incidence of POA. There was a correlation between age and agitation severity as measured by Pearson analysis (correlation coefficient=0.75). Females showed a lower incidence of postoperative agitation; the study included 77 males and 59 females, which represent 87% versus 22% in males and females respectively. Chi-square test revealed an association between gender and agitation with the correlation coefficient 0.725, P<0.0001 (Table 2). In the total sample size, there were 95 nonsmokers, and 42 smokers, the incidence of agitation was found to be 88% among smokers versus 44% for nonsmokers P<0.0001. There was no statistical difference in pain score between the two studied groups, pain scores of 4.7±1.2 versus 4.4±1.3 in pre-medication and the control group respectively, P=0.207 however, there was a correlation between postoperative pain, and agitation (Table 2). There was no significant correlation between agitation and either postoperative O2 saturation or BIS (Table 2).

Table 2: Correlation coefficients between variables and total agitation.

<table>
<thead>
<tr>
<th>Test</th>
<th>Smoking index</th>
<th>SPO2</th>
<th>Pain</th>
<th>BIS</th>
<th>age</th>
<th>gender</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spearman coefficient of determination</td>
<td>0.578</td>
<td>0.1</td>
<td>0.25</td>
<td>0.01</td>
<td>0.43</td>
<td></td>
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<tr>
<td>ANCOVA R²</td>
<td>0.3</td>
<td>0.08</td>
<td>0.2</td>
<td>0.01</td>
<td>0.34</td>
<td></td>
</tr>
<tr>
<td>Chi square</td>
<td>0.671</td>
<td>0.06</td>
<td>0.389</td>
<td>0.01</td>
<td>0.75</td>
<td>0.72</td>
</tr>
<tr>
<td>P value</td>
<td>&lt;0.0001</td>
<td>0.07</td>
<td>0.001</td>
<td>0.333</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Discussion

The main finding of the present study is that premedication with atropine and Phenergan increased the occurrence and severity of postoperative agitation in adults following functional endoscopic sinus surgery. Postoperative agitation in adults has not been adequately studied; few studies have reported the incidence postoperative agitation in adult patients following nasal surgery [9,10]. A plenty of medications have been shown to increase delirium and excitations, [11] anticholinergics and antihistaminic are of particular concern as these drugs are routinely used in the pre-anesthetic medications [12]. The results of the present study reveal that pre-medication with both Atropine (an anticholinergic) and Phenergan (an antihistamine) substantially increase the occurrence of postoperative agitation following FESS. Because both Atropine and Phenergan cross the blood brain barrier, they may interact to cause central excitation that is noted as...
postoperative agitation. These agitations could be minimized by the use of a quaternary ammonium anticholinergic like glycopyrrolate as an Atropine-substitute. Some postoperative events might provoke agitation, postoperative pain [13] and hypoxia [14], are major confounders that may cause or result from agitation. While hypoxia and pain can cause agitation, an agitated patient can remove his O2 mask or injure himself resulting in more hypoxia and pain. In our study, because none of the patients showed are duction in recovery oxygen saturation, there was no correlation between agitation and the recovery oxygen saturation; this would exclude hypoxia as a confounding factor for postoperative agitation. In the present study, there was a correlation between agitation and postoperative pain, indicating that pain was a contributing factor in the development of agitation. In spite of that, pain cannot be considered the only cause of POA because, in the present study, the postoperative pain was equal in both groups while premedication group displayed a significantly higher agitation. In addition, the correlation coefficient between pain and agitation was found to be 0.25, and this denotes that pain can explain 25% only of the agitation. Consistent with this, Weldon, et al. reported that emergence agitation could occur even after adequate pain treatment or after procedures that are not associated with pain [15].

Postoperative agitation in adults was reported mainly in geriatric population [16,17]. Surprisingly more frequent and more severe agitation were detected among 18 to 30 years old group of patients. We expected the reverse to occur as delirium was almost recorded in geriatric patients [18,19]. It is apparent that this type of POA represents a separate entity than those noticed for geriatric populations. Although both POA and delirium are manifested by patient confusion and excitement, it appears that the etiology and predisposing factors are different. POA in our study mimics those displayed by children anesthetized with sevoflurane rather than that shown in geriatric populations. Agitation was more prominent among smokers, the matter that raises the question whether the increased agitation is due to nicotine effects or nicotine withdrawal? Nicotine withdrawal during the fasting period may cause restlessness and irritability. Nicotine withdrawal has been shown to be associated with a negative emotional state, including anxiety and the perception of increased stress [20,21]. Withdrawal symptoms include irritability, depressed mood, restlessness, anxiety, difficulty concentrating and insomnia [20]. It is possible that nicotine abstinence during the fasting period (8 hours), duration of surgery average (2) hours have accounted for the increased agitation among smokers. Further studies are required to clarify this issue. Agitation was more frequent and severe among males than females; one possible explanation is the hormonal difference between both sexes. Interestingly, testosterone has been shown to decline with advancing age [22]. So it is possible that higher testosterone level in young males has contributed to increased agitation in this group. Further studies are clearly required to address this issue. Intraoperative awareness has not been detected during our study as measured by the bispectral index. Therefore, in our protocol, we can exclude awareness as a risk factor for postoperative agitation. Nevertheless, the accuracy of BIS as a tool to measure intraoperative awareness is controversial [23-28]. It remains possible that some degree of awareness has occurred during hypotension where inhalational anesthetic concentration was reduced. There are no data on the effect of hypotension on BIS value. Furthermore, it is not clear if this level of hypotension has suppressed brain electrical activity to yield a significant reduction in BIS recording. It remains possible that an unknown effect of hypotension has decreased BIS value. This reduction was counteracted by some degree of awareness during hypotension when the inhalational anesthetic concentration was reduced, so the net value was equal. Emergence agitation is costly in several ways, in terms of morbidity, in human resources and on the financial level [29]. It is well-known that personnel costs account for the majority of PACU costs [30,31]. The agitated patient requires 4-6 recovery nurses to control his movement and apply restraints, take care of monitoring. In our study agitated patients removed oxygen masks, IV cannulas, nasal pack, pulse oximeter probes several times before being quiet, the matter that shifted attention to this particular patient, rendered many recovery staff busy at the expense of the other patients. We acknowledge that our study has some limitations. First, we did not study the effect of hypotension on agitation because all patients were subjected to induced hypotension to reduce bleeding and improve the surgical field. Second, the role of the nasal pack in the development of POA was not investigated. Further studies are needed to answer these questions.

**Conclusion**

a. Premedication with Atropine and Phenergan increase the incidence of postoperative agitation. Avoiding premedication or the use of their substitutes might reduce the incidence of postoperative agitation.
b. Postoperative pain, smoking, young age, male gender were identified as risk factors for postoperative agitations following FESS.

- **Author Contribution:** Hazem E. Elsersy: Idea, experimental design, patient recruitment, randomization, data analysis, manuscript writing. Ahmed A A: experimental design, data analysis.

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- **Conflict of interest:** none declared

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