Sociodemographic Status of Propofol and Dexmedetomidine as an Agent of Conscious Sedation in Patients Undergoing Tympanoplasty

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Abstract

Objective: In this study our main goal is to assess sociodemographic characteristics of propofol and dexmedetomidine as an agent of conscious sedation in patients undergoing tympanoplasty. Method: This Randomised prospective clinical trial study was carried out with the patients who underwent tympanoplasty conducted at ENT operation theatre in BSMMU, Dhaka according to inclusion & exclusion criteria from February 2016 to February 2018. ASA I-II patients aged (18-45) years undergoing tympanoplasty were enrolled. Patients were randomized as Group I and Group II by computer-gerenated randomization where there was 30 patients in each group (n=30). Results: Both groups were almost similar in respect to age, weight, height, BMI. Mean age of the patient group-I was 30.93 ± 7.11 and 29.03 ± 7.36 in group-II and among them maximum age was 50 years and minimum age was 19 years in group-I and 44,14 years in group-II. Mean weight of the patient group-I was 58.43 ± 6.88 and 57.50± 5.52 group-II and among them maximum weight was 72 kg and minimum weight was 45 kg in group-I and 67,49 years in group-II. Mean height of the patient group-I was 1.60 ±0.06 and 1.59 ± 0.06 group-II and among them maximum height was 1.73 and minimum height was 1.49 in group-I and 1.79,1.49 meter in group-II. Mean BMI of the patient group-I was 22.72 ± 2.60 and 22.52 ± 2.18 group-II. Demographic data between the two groups were not statistically significant. During the study, a computer-generated randomization table was used to allocate the patients into 2 equal groups, 30 in each where Group I received dexmedetomidine and Group II received propofol. Observations were made during perioperative period for conscious sedation effectiveness on haemodynamics SBP, MAP, DBP, HR, SpO2 on different time intervals, surgical field bleeding status, patient satisfaction, surgeon satisfaction, intraoperative rescue sedation, intraoperative rescue analgesics, postoperative rescue analgesics, time to achieve target Ramsay sedation scale (RSS) after induction, intra and postoperative pain intensity level, and adverse effects. Demographic data and different parameter were described as mean with standard deviation in different tables and graphical presentation on different types of charts. Conclusion: From our results we can conclude that, conscious sedation with dexmedetomidine with local anaesthetic lignocaine infiltration provides better outcome of conscious sedation in terms of patient’s satisfaction, surgeon’s satisfaction, reduce per operative surgical site bleeding for patients undergoing tympanoplasty. Keywords: Propofol, Dexmedetomidine, haemodynamics. Tympanoplasty.

INTRODUCTION

There are various types of surgeries done in ears including external, middle and internal ear either individually or simultaneously. This procedure done under different types of anaesthetic technique like general anaesthesia, local infiltration, local infiltration with sedation, regional blocks and or combination all above procedures. Each of technique has its own advantages and disadvantages.
Using local anaesthesia only comparison to general anaesthesia in terms of patient recovery, patient turnover, and avoidance of intubation and elimination of possibility of laryngotracheal complication is as well as removal of cardiovascular, respiratory, renal, and neurological complications. Local anaesthesia also reduce intraoperative bleeding and helps perioperative hearing assessment, vertigo detection required during surgery. It also prevents potentially lengthy and distressive emergence from general anaesthesia.

Propofol is a newer drug has been used for conscious sedation have narrow therapeutic index, risk of progression to deep sedation. It has no analgesic effect and must be used with adequate pain relief. Propofol causes hypotension, apnoea required intervention during conscious sedation [1]. It also causes hypoxia, vomiting, deep sedation, coughing, agitation, airway obstruction, desaturation, higher recall [2].

Therefore, suitable drugs and adequate doses for sedation have been debated and variety of drugs used around the world used and consequently search for appropriate newer drugs continues. The alpha 2 agonist dexmedetomidine now a day’s widely used for its sedative analgesic and sympatholytic properties in the perioperative and critical care. It was approved by FDA 1999 for used as sedative agents [3]. It has shorter half-life and eight fold greater sensitivity and selectivity for the receptor than clonidine [4]. It also attenuates the stress response to surgery, provides opioid sparing effect, stable haemodynamics [5]. It shows that intra operative administration of dexmedetomidine maintain haemodynamic stability. It prevents post-operative nausea, vomiting, shivering and potential benefits of cardio, neuro and renal protection [6]. It can be used in perioperative period as an analgesic adjunct. Lower heart rate and mean arterial pressure provides better operative field for microscopic surgery [7]. It decreases bleeding when bloodless surgical field required. Dexmedetomidine used significantly less tramadol in FESS and septoplasty [8]. It is better drug for conscious sedation with better haemodynamic stability; reduce analgesic requirements fentanyl about 44% [9].

Significantly higher rates of effective sedation in dexmedetomidine compared with midazolam [10]. It has calmer, cooperative, higher satisfaction score without compromising airway. It is safe to use in impending respiratory failure [11]. Dexmedetomidine has been used for various surgical procedures, preserved muscle tone and spontaneous ventilation and awaken by external stimulus [12]. EEG study demonstrated sedative effects of dexmedetomidine mimics second stage of non REM sleep [13]. In USA dexmedetomidine approved for sedation of non-intubated patient or during surgical procedure. A Conchrane review 2009 examined benefits of alpha 2 agonist in obtundng the perioperative stress induced sympathetic activity [14]. In CNS dexmedetomidine stimulates brainstem reduce heart rate and decrease blood pressure [15]. It has selective alpha 2 agonist with properties of analgesia, sympatholysis and can titrate sedation without respiratory depression. It reduces opioid requirements and stress response to surgery ensuring a stable haemodynamics [16]. It is increasingly used for various surgical procedures. The use of dexmedetomidine in other ENT surgeries like FESS, septoplasty, thyroplasty under conscious sedation has also documented [17]. It can be safely and effectively used for surgeries under conscious sedation. At low or moderate doses with slow rates of infusion of dexmedetomidine alpha 2 agonist effects are observed but no alpha 1 effect. It has been used as sedatives and analgesic as single agent in many procedures [18].

In our country few articles published on dexmedetomidine shows that dexmedetomidine infusion reduces the NT-Pro BNP level has better cardiac outcome for patients undergoing ORIF for fracture shaft of femur. But no such study yet did on tympanoplasty at Bangladesh [19]. We assume that conscious sedation with dexmedetomidine may be a better alternative to other sedative drugs for patient undergoing tympanoplasty.

OBJECTIVE
General objective
Assess sociodemographic characteristics of propofol and dexmedetomidine as an agent of conscious sedation in patients undergoing tympanoplasty.

Specific objective
- To compare intra operative haemodynamics. (SBP, DBP, MAP, SPO2, HR).
- To compare operative field bleeding by bleeding scale.
- To compare time to achieve adequate sedation by Ramsay sedation scale.
- To compare time of full recovery from sedation by modified Aldrete score.

METHODOLOGY
Study type
- This was an Randomised prospective clinical trial.

Study place and period
This study was carried out with the patients who underwent tympanoplasty conducted at ENT operation theatre in BSMMU, Dhaka according to inclusion & exclusion criteria from February 2016 to February 2018

Sampling method
- The sample was collected by computer generated random sampling.
Study population
- Total 30 patients were included in this study.

Selection criteria
Inclusion criteria
- All Patients (both male and female) underwent tympanoplasty.
- Aged between 18-45 years of age.
- ASA physical status I and II

Exclusion criteria
- Patient (both male & female) refused to be included in the study.
- COPD, asthma, Cardiac, renal, hepatic dysfunction or disease.
- History of difficult intubation.
- Coagulopathy or coagulation disorder.
- Obesity (20% of ideal body weight).
- History of drug allergy.
- Patients requiring endotracheal intubation.
- Sleep apnoea.

Procedure of data collectio
This randomized clinical trial took place at BSMMU in ENT operation theatre under Department of Anaesthesia, Anægæsia and Intensive care medicine after approval by Institutional Review Board (IRB). Patients were randomized as Group I and Group II by computer-generated randomization where there was 30 patients in each group (n=30). Patients were interviewed for detailed medical and drug history and underwent a physical examination before the surgery to verify whether fulfill inclusions criteria. Before the surgery, patients were instructed clearly about anaesthetic technique, NRS pain scale.

The group I received dexmedetomidine and group II received propofol. After taking the patient on the operation table, a monitoring device was attached and baseline haemodynamics (SBP, DBP, MAP, SpO2, HR) were noted. An intravenous cannula was inserted for giving intravenous fluids and drugs. All patients were monitored with an automated noninvasive blood pressure device, pulse oximetry, and an electrocardiogram. Drugs were prepared and two 50 ml syringes were labeled as loading and maintainence syringe for each patient.

Patients in Group I were get injection dexmedetomidine loading dose 1µgm/kg over 10 minute followed by continuous infusion 0.4 µgm/kg/hr. Patients in the Group II were get injection propofol loading dose 75 µgm/kg intravenously over 10 minute than 50µgm/kg/min continuous infusion was started.

Patients were observed for the depth of sedation throughout the period of surgery and sedation both intraoperatively and postoperatively. Light sedation was assumed from the observation of

Statistical analysis
All relevant collected data was compiled on a master data sheet first. Then organized. Statistical analyses were carried out by using the Statistical Package for Social Sciences version 16.0 for Windows (SPSS Inc., Chicago, Illinois, USA). The mean values were calculated for continuous variables. The quantitative observations will be indicated by frequencies and percentages. Chi-Square test was used to analyze the categorical variables like sex, ASA status and surgical satisfaction, patient satisfaction which was shown with cross tabulation. Un paired t-test was used for continuous variables like systolic blood pressure (SBP), diastolic blood pressure (DBP), mean arterial pressure (MAP), heart rate (HR), percent saturation of oxygen (SpO2) at different interval. Unpaired t-test was also be used for age, weight, height, duration of surgery. P values <0.05 was considered as statistically significant.

Results
Demographic data
Both groups were almost similar in respect to age, weight, height, BMI. Mean age of the patient group-I was 30.93 ± 7.11 and 29.03 ± 7.36 in group-II and among them maximum age was 50 years and minimum age was 19 years in group-I and 44, 14 years in group-II.

Mean weight of the patient group-I was 58.43 ± 6.88 and 57.50± 5.52 group-II and among them maximum weight was 72 kg and minimum weight was 45 kg in group-I and 67.49 years in group-II. Mean height of the patient group-I was 1.60 ±0.06 and 1.59 ± 0.06 group-II and among them maximum height was 1.73 and minimum height was 1.49 in group-I and 1.79, 1.49 meter in group-II. Mean BMI of the patient group-I was 22.72 ± 2.60 and 22.52 ± 2.18 group-II.

Demographic data between the two groups were not statistically significant.
Table-I: Demographic data and clinical data (n = 30 each group)

| Parameter             | Groups                  | P value  |
|-----------------------|-------------------------|----------|
| (n = 30 each group)   | Group I                 | Group II |          |
| Age                   | 30.93 ± 7.11            | 29.03 ± 7.36 | 0.31 "    |
| Range                 | 19.45                   | 14.44    |          |
| Weight                | 58.43 ± 6.88            | 57.50 ± 5.52 | 0.56 "    |
| Range                 | 45.72                   | 49.67    |          |
| Height                | 1.60 ± 0.06             | 1.59 ± 0.06 | 0.72 "    |
| Range                 | 1.49 ± 1.73             | 1.49 ± 1.79 |          |
| BMI                   | 22.72 ± 2.60            | 22.52 ± 2.18 | 0.75 "    |
| ASA grade             |                         |          |          |
| I                     | 28(93.3%)               | 26(86.7%) | 0.38 "    |
| II                    | 2(6.7%)                 | 4(13.3%) |          |
| Marital status        |                         |          |          |
| Married               | 24(80%)                 | 17(56.7%) | 0.09 "    |
| Unmarried             | 6(20%)                  | 13(43.3%) |          |

Significant ", non-significant "ns

Values were expressed as number and percentages.

"p was derived from chi square test, "p was derived from unpaired t test.

**Perioperative haemodynamics**

**Mean arterial blood pressure**

Base line mean± SD value of MAP in Group I was 84.91 ± 3.49 mm of Hg. DBP during different evaluation period varied from 82.87 ± 2.91 mm of Hg to 83.35 ± 3.17 mm of Hg. On the other hand base line MAP of Group II was 83.12 ± 4.20 mm of Hg and during different evaluation period ranged from 82.77 ± 4.12 mm of Hg to 85.42 ± 4.32 mm of Hg. The changes were similar most of the observation and showed no significant differences between groups.
Fig-II: Line diagram showing comparison of MAP of two groups in different time intervals throughout the perioperative period. (Mean)

Time to achieve target RSS

Time to achieve target RSS after induction. Among group I, 24(80%) patient achieves target sedation (RSS = 3) within (8-10) minutes and 3(10%) within (5-8) minutes and 3(10%) patients’ needs more than 10 minutes to achieve target RSS. On the contrary among group II, 25(83.3%) patient achieves target sedation (RSS = 3) within (1-5) minutes and 4(13.3%) within (5-8) minutes and 1(3.3%) patients’ needs more than 10 minutes to achieve target RSS. Study revealed Group II found statistically significant (P<0.05).

Table-II: Ramsay sedation score (n = 30 each group)

| Groups     | (1-5) minute | (5-8) minute | (8-10) minute | Above10 minute | *pvalue |
|------------|--------------|--------------|---------------|----------------|---------|
| Group I    | 0%           | 10%(3)       | 80%(24)       | 10%(3)         | 0.01**  |
| Group II   | 83.3%(25)    | 13.3%(4)     | 0%            | 3.3%(1)        |         |

Significant *

Values were expressed as number and percentages. 
*p was derived from chi square test.

Intraoperative pain intensity

Intraoperative pain intensity score at different grade in intraoperative period. Among group I, 23 (76.7%) patients have no pain, 4(13.3%) have mild pain 3(10%) have moderate pain with the anaesthesia technique. On the contrary 21(70%) have no pain, 3 (10%) have mild pain and 6(20%) have moderate pain with the anaesthesia technique in group II respectively. Study revealed Group II found statistically significant (P<0.05).

Table-III: Pain intensity (n= 30 each group)

| Parameters    | Groups     | No pain | Mild pain | Moderate pain | Severe pain | *pvalue |
|---------------|------------|---------|-----------|---------------|-------------|---------|
| Intraoperative period | Group I    | 76.7%(23) | 13.3%(4)  | 10%(3)        | 0%          |         |
|                | Group II   | 70%(21)  | 10%(3)    | 20%(6)        | 0%          |         |
| Postoperative period | Group I    | 63.3%(19) | 20%(6)    | 16.6%(5)      | 0%          |         |
|                | Group II   | 66.7%(20) | 30%(9)    | 3.3%(1)       | 0%          | 0.2*** |

Significant *, nonsignificant **

Values were expressed as number and percentages. 
*p was derived from chi square test.
Intra operative rescue sedation

Rescue sedation during intraoperative period. Among group I, 25(83.3%) needs no rescue sedation, and only 5 (16.7 %) patients required rescue sedation with the anaesthesia technique. On the contrary 24(80%) patients not required rescue sedation and only 6(20%) patients required rescue sedation in group II respectively. Study revealed found statistically not significant.

Surgical field bleeding

Surgical field assessed by bleeding score. Bleeding score is obtained during intraoperative period. Among group I, we found 19 (66.7%) had score 0, which was considered no bleeding and the rest 11(33.3%) scored 1, was considered as mild bleeding with the anaesthesia technique required no aspiration. On the contrary only 9(30%) had score 0, considered as no bleeding and the rest and most 17(56.7%) patients had score 1, was considered as mild bleeding with the anesthesia technique required frequent aspiration and 1 (3.3%) patient had score 4 was considered moderate bleeding in group II respectively. No group had scored 5 which were considered as severe bleeding. Study revealed Group I found statistically significant (P<0.05).

| Groups | No bleeding(0) | Mild bleeding (1) | Mild bleeding (2) | Mild bleeding (3) | Moderate bleeding (4) | Severe bleeding (5) | P value |
|--------|----------------|------------------|------------------|------------------|----------------------|---------------------|---------|
| Group I | 66.7%(19)      | 33.3%(11)        | 0%               | 0%               | 0%                   | 0%                  | 0.04*   |
| Group II | 30%(9)         | 56.7%(17)        | 3.3%(1)          | 6.7%(2)          | 3.3%(1)              | 0%                  |         |

Table-V: Satisfaction score (n = 30 each group)

| Parameters       | Groups         | Very dissatisfied | Dissatisfied | Neutral | Satisfied | Very satisfied | P value |
|------------------|----------------|-------------------|--------------|---------|-----------|----------------|---------|
| Patient satisfaction | Group I       | 0%                | 3.3%(1)      | 6.7%(2) | 13.3%(4)  | 80%(23)        | 0.01*   |
| Surgeon satisfaction | Group II     | 0%                | 0%           | 6.7%(2) | 3.3%(2)   | 90%(26)        |         |
|                  | Group I       | 0%                | 0%           | 13.3%(4) | 73.3%(22) | 13.3%(4)       | 0.01*   |

Aldrete score

Patient satisfaction, surgeon satisfaction, time of achievement of modified Aldrete score was statistically significant.
In this present study, Rescue analgesia during intraoperative period. Among group I, we found 23(76.7%) patients' needs no rescue analgesia, and only 7(23.3%) patients required rescue analgesia with the anaesthesia technique. On the contrary 10(33.3%) patients required rescue analgesics and 20(66.7%) patients not required rescue analgesics in group II respectively. Study revealed Group II found statistically significant (P<0.05) consistent with the result of [7].

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In this study, it was observed that percentage of patient in group I achieve target VAS value were found significantly lower in dexmedetomidine group compared to midazolam as the target VAS value were found significantly higher VAS in midazolam group compared to propofol. Our study also corresponds to and it was statistically significant.

It was observed that Among group I, 25(83.3%) needs no rescue sedation, and only 5 (16.7%) patients required rescue sedation with the anaesthesia technique. On the contrary 24(80%) patients not required rescue sedation and only 6(20%) patients required rescue sedation in group II respectively. The total number of rescue doses of sedatives was lesser in dexmedetomidine group consistent with the findings of [7].

In this current study, it was observed that bleeding score was less in dexmedetomidine group and provides better surgical condition in comparison to propofol. Among group I, we found 19 (66.7%) had score 0 on the contrary only 9(30%) had score 0 in group II. Clonidine reduces blood loss and provide better surgical condition in comparison to midazolam. Clonidine and Dexmedetomidine both are found effective in reducing bleeding in ENT surgeries [25].

In conclusion this prospective randomization study demonstrated that comparing the

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dexmedetomidine and propofol for conscious sedation in tympanoplasty we found that dexmedetomidine have better propofol for sedation in the emergency department. Academic Medicine. 1996 Mar;3(3):234-8.

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