Comparison of Sedation with Dexmedetomidine Alfentanil Versus Ketamine-Alfentanil in Patients Undergoing Closed Reduction of Nasal Fractures

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Abstract

Objectives: To compare the sedative effects of dexmedetomidine alfentanil versus ketamine-alfentanil in patients undergoing closed reduction of nasal fractures on the basis of intraoperative hemodynamic changes, satisfaction of patients and surgeons, and the adverse effects.

Methods: Sixty patients with ASA class 1 or 2 were randomized to either of two groups, a dexmedetomidine alfentanil group (DA group; n = 30) or a ketamine-alfentanil group (KA group; n = 30). Hemodynamic parameters, oxygenation status, adverse events, the satisfaction of patients and surgeons, and postoperative pain scores by visual analog scale (VAS) were recorded at specific time intervals during the trial.

Results: Systolic blood pressure was significantly lower in the DA group than in the KA group from T1 min to T15 min. The duration of the recovery ward stay was longer in the DA group; however, two groups were similar in terms of total anesthesia time and awakening time. Likewise, two groups were similar in terms of the patient and surgeon’s satisfaction, pain scores, and the occurrence of adverse effects.

Conclusions: Both sedation methods were safely performed, and dexmedetomidine-alfentanil is as effective as ketamine-alfentanil in patients undergoing short-term operations such as nasal fracture corrections.

Keywords: Dexmedetomidine, Ketamine, Alfentanil, Nasal Fracture

1. Background

Nasal bone fractures are the most common type of facial fractures (1). One treatment for this type of fracture is a closed reduction that can be done under local or general anesthesia (2). General anesthesia provides better patient satisfaction with anesthesia, appearance, and function of the nose and reduces the corrective surgeries such as septoplasty, septrhinoplasty, and rhinoplasty (3). However, it may not be an option because of medical risk factors, emergent need for intervention, or operating room availability. Intravenous sedation may be the most controllable way of determining the exact level of ideal sedation individually needed for short-term operations such as nasal manipulations (4, 5).

Dexmedetomidine is a highly selective α₂ agonist that reduces peripheral norepinephrine secretion by stimulating inhibitory prejunctional α₂ adrenoreceptors. The half-life of dexmedetomidine is 2.3 hours and its distribution half-life is less than 5 minutes; so, its clinical effects are very rapid (6-8). Also, it assists hemodynamic stability by its sympatholytic action, which has been used as a safe adjuvant during anesthesia in many clinical applications (9-12). Dexmedetomidine has been used as a sedative and hypnotic in procedures that do not need tracheal intubation. Since dexmedetomidine has no negative effect on respiratory rate or depth than other sedatives, it has been proven that it is usable in airway-related procedures (13, 14). However, ketamine is a dissociative anesthetic that plays a significant role in analgesia and sedation for outpatient surgeries. It has not only sedative-hypnotic properties resulting in a light dissociative sleep but also has potent analgesic properties (15, 16).

Closed reduction of nasal fractures is a short-term operation that needs awakening closely after anesthetic induction. Therefore, the opioids administered during the
operation should be short-acting to provide a rapid awakening from anesthesia. Alfentanil is a potent, rapid, and short-acting analog of fentanyl, which appears to have some significant advantages during outpatient surgeries (17).

2. Objectives

The aim of this study was to compare the sedative effects of dexmedetomidine alfentanil versus ketamine-alfentanil in patients undergoing closed reduction of nasal fractures.

3. Methods

This randomized, double-blind clinical trial (code: IRCT20191127045530N1) was approved by the Ethics Committee of the Ahvaz Jundishapur University of Medical Sciences (code: IR.AJUMS.REC.1398.624) and was conducted in Imam Khomeini Hospital, Ahvaz, Iran from Jan to February 2020. The advantages and disadvantages of sedation were explained to the patients before the operation.

The patients were randomly divided into two groups, including the dexmedetomidine alfentanil group (DA group; n = 30) and the ketamine-alfentanil group (KA group; n = 30). Patients with ASA class I or II, aged 15 to 40, were scheduled to reduce nasal bone fracture. Patients under 15 and over 40 years, patients with a history of analgesic consumption within 24 hours before the surgery, uncontrollable hypertension or heart diseases, coagulopathy, a history of convulsion, and neurologic disorders were excluded from the study.

Blood pressure, heart rate, and peripheral oxygen saturation (SpO₂) were continuously monitored in patients entered the operating room every 5 min. At least 5 min prior to the operation, all patients received supplementary 100% oxygen at a rate of 5 - 6 lit/min through a face mask.

Both groups received 0.05 mg/kg midazolam (Chemidarou, Iran) and 0.01 mg/kg atropine (Caspian, Iran) intravenously before anesthetic induction. Then, in the dexmedetomidine-alfentanil (DA) group, dexmedetomidine (Exir, Iran) was administered at a rate of 1 µg/kg over 10 min in combination with 8 µg/kg alfentanil (Darou Pakhsh, Iran) to give a sedation level of 5 - 6 according to Ramsay sedation scale (Table 1). In the ketamine-alfentanil (KA) group, sedation was induced with bolus injection of ketamine 1mg/kg (Rotexmedica, Germany), 8 µg/kg alfentanil to keep the same level of sedation. Discomfort and agitation were considered indicators of inadequate sedation and were treated with i.v. Propofol 20 mg (Fresenius Kabi, Austria).

In both groups, changes in sBP, dBP, MAP, HR, and SpO₂ were recorded by staff members who were blinded to the patients’ group allocation at specific time intervals. Anesthesia time, awakening time (based on Ramsay sedation score: 3), and the recovery ward stay duration (based on Modified Alderete score: 9) were recorded. After surgery, satisfaction of the patients and surgeon was measured (7-point Likert-like verbal rating scale), and the pain was evaluated using the visual analogue scale (VAS, 0: no pain-10 cm: worst pain) at specific intervals until six hours.

All variables were evaluated by an investigator who was blinded to the patient group. SPSS 26 software was used for statistical analysis. All values were expressed as mean (SD), median (range), or the number of patients (%). Student t-test was used to compare continuous variables, and repeated measures analysis of variance (ANOVA) was used to analyze hemodynamic changes over time. A chi-square test was used to analyze categorical variables. P values of less than 0.05 were considered statistically significant.

4. Results

There was no statistically significant difference between the groups in age, sex, and weight of the patients (Table 2). Two groups were similar in terms of the patient and surgeon’s satisfaction and also no significant differences in the occurrence of nausea, agitation, and lethargy (Table 3). The postoperative pain was assessed using the Visual Analogue scale (VAS) before induction, in the recovery ward, 2, and 6 hours postoperatively. The results showed that there was no significant difference between the two groups (Table 4). The duration of the recovery ward stay was longer in the DA group (36.83 (8.03) min vs. 30.00 (7.76) min, P < 0.001); however, the two groups were similar in terms of total anesthesia time and awakening time (Table 5).

| Table 1. Ramsay Sedation Scale (18) |
|-----------------------------------|
| Score | Details |
| 1     | Patient is anxious or agitated |
| 2     | The patient is cooperative, oriented, and tranquil |
| 3     | Patient responds to commands only |
| 4     | Patient exhibits brisk response to light glabellar tap or loud auditory stimulus |
| 5     | Patient exhibits a sluggish response to light glabellar tap or loud auditory stimulus |
| 6     | Patient exhibits no response |

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Regarding intraoperative hemodynamic changes, the DA group had a lower systolic BP, MAP, and HR compared to the KA group, but only systolic blood pressure changes from T1 min to T15 min were significantly different. (P < 0.01) (Figure 1).

No episodes of desaturation (SpO₂ < 90%) were noted during this study and there was no significant difference in SpO₂ between both groups. (Figure 2)
Ketamin 
Dex 
Ketamin 
Dex 
Ketamin 
Dex 
Ketamin 
Dex 
Time 
Time 
Time
Time
Heart Rate (Rate/min) ... 
65 
T0 T1 T5 T15 T2h T6h Trecov T0 T1 T5 
T0 T1 T5 T15 T2h T6h Trecov T0 T1 T5 T15 T2h T6hTrecov 
T15 T2h T6h Trecov

Figure 1. Changes in hemodynamic variables. Data are shown as mean. DA: dexmedetomidine-alfentanil, KA: ketamine-alfentanil. T0: baseline, T1, T5, and T15: 1, 5, and 15 min after drug infusion in both groups, Trecov: at the recovery ward, T2h, and T6h: 2 and 6 hours after infusion.

Figure 2. Changes in peripheral oxygen saturation (SpO2). Data are shown as mean DA: dexmedetomidine-alfentanil, KA: ketamine-alfentanil. T0: baseline, T1, T5, and T15: 1, 5, and 15 min after drug infusion in both groups, Trecov: at the recovery ward, T2h, and T6h: 2 and 6 hours after infusion.

5. Discussion

Sympatholytic property of dexmedetomidine provided lower systolic blood pressure during T1 min to T15 min of operation. Our finding was in accordance with studies that found a significant reduction in hemodynamic status using dexmedetomidine (19, 20). Also, Jae-Wook et al. (21) demonstrated that dexmedetomidine has sympatholytic effects and it is used for sedative, anxiolytic, and analgesic purposes as an adjunct to several anesthetics. So, these complex properties of dexmedetomidine led patients to experience less pain during the postoperative period in their study.

Kim et al. (22) stated that pre-operative administration of dexmedetomidine in a closed reduction of nasal bone fractures reduces agitation severity and shortens the duration of agitation and also provided more stable maintenance of anesthesia with less movement during the operation.

Our results confirmed the findings of Kim et al. (23) who conducted a randomized controlled trial of 100 nasal surgeries (the dexmedetomidine group received dexmedetomidine infusion, while the control group received normal saline as the placebo). Intra-operative continuous dexmedetomidine infusion resulted in more...
stable hemodynamics during extubation and enhanced patient-reported global quality of recovery 24 h after surgery.

The results of this study showed that two groups were similar in terms of the patient and surgeon’s satisfaction and postoperative pain scores, and also, there were no significant differences in the occurrence of adverse effects. In a similar study, Lee et al. (24) found that monitored anesthesia care with dexmedetomidine resulted in comparable satisfaction in the patients, nurses, and surgeons compared to general anesthesia. The occurrence of complications and pain scores were also similar.

One of the most common problems in the recovery ward is delayed recovery and longer discharge time of patients receiving pre-operative or intra-operative dexmedetomidine infusion. (19, 21, 22) In our research, the duration of the recovery ward stay was significantly longer in the DA group than in the KA group. However, two groups were similar in terms of total anesthesia time and awakening time.

5.1. Conclusions

This study was conducted to compare dexmedetomidine-alfentanil versus ketamine-alfentanil for sedation of patients undergoing closed reduction of a nasal bone fracture. Both sedation methods were performed without the incidence of any serious side effects, and two groups were similar in terms of the patient and surgeon’s satisfaction, pain scores and occurrence of complications. Therefore, dexmedetomidine-alfentanil is considered to be a good choice for sedation during nasal fracture corrections.

Footnotes

Authors' Contribution: None declared by author.

Clinical Trial Registration Code: The clinical trial registration code was IRCT20191127045530N1.

Conflict of Interests: No conflict of interest was declared.

Ethical Approval: The ethical approval code was IR.AJUMS.REC.1398.624.

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