ISSN (Print): 2209-2870 ISSN (Online): 2209-2862





International Journal of Medical Science and Current Research (IJMSCR)

Available online at: www.ijmscr.com Volume 6, Issue 1, Page No: 18-30

January-February 2023

# A Randomized Study Of Midazolam Versus Dexmedetomidine For Premedication In Pediatric Age Group Via Intranasal Mucosal Atomization Device

<sup>1</sup>Dr. Ashish Tyagi, <sup>2</sup>Dr. Malini Mehta, <sup>3</sup>Dr. Garima Sinha, <sup>4</sup>Dr. Shailaja Tyagi, <sup>5</sup>Dr. Rashmi Kumari <sup>3</sup>Assistant Professor, <sup>2</sup>HOD and Professor, <sup>4</sup>Consultant, <sup>5</sup>Junior Resident,

<sup>3</sup>Department of Anaesthesia and Critical care, GIMS Hospital, Greater Noida, Uttar Pradesh, India <sup>5</sup>Department of Critical care, <sup>1</sup>Department of Anaesthesiology,

<sup>1,5</sup>School Of Medical Sciences And Research, Sharda University, Greater Noida. Uttar Pradesh, India <sup>2</sup>Department of Emergency Medicine, Shrimati Bhikiben Kanjibhai Shah Medical Institute and Research Centre, Sumandeep Vidyapeeth (An Institute Deemed to be university), Piparia, Vadodara, Gujarat, India <sup>4</sup>Department of Pathology, Yatharth Super specialty Hospital, Noida Extension, Greater Noida, Uttar Pradesh, India

## \*Corresponding Author: Dr. Ashish Tyagi

Assistant Professor, Department of Anaesthesiology, School of Medical Sciences And Research, Sharda University, Greater Noida. Uttar Pradesh, India

Type of Publication: Original Research Paper

Conflicts of Interest: Nil

#### **Abstract**

### **Introduction:**

Many drugs have been tried for premedication in children<sup>[1]</sup> amongst which midazolam and dexmedetomidine are commonly used and are reportedly safe and effective for usage during both separation as well as induction of anaesthesia. Hence, we planned to carry out a study of midazolam versus dexmedetomidine for premedication in children via intranasal Mucosal Atomization Device.

### Aim of the study:

To compare the efficacy of midazolam and dexmedetomidine as premedication in pediatric age group of 2 to 6 years of age via intranasal Mucosal Atomization Device.

### Materials and methods:

This randomized study was done at Dhiraj hospital, Piparia, Vadodara, Gujarat on 60 children belonging to American Society of Anaesthesiologists (ASA) physical status I or II of age 2-6 years of either gender undergoing elective surgeries under general anaesthesia. Children were separated into: Group M - Midazolam 0.4 mg/kg (preservative free) Group D - Dexmedetomidine 2µg/kg. Drug was administered 30 minutes prior to surgery and following parameters were assessed: Acceptance of drug, Sedation Score, behaviour during parental separation and mask acceptance.

### **Results:**

Dexmedetomidine was statistically significantly better in aspect of drug acceptance, sedation after 30 minutes of drug administration, parental separation and mask acceptance than midazolam.

## **Conclusion:**

Intranasal dexmedetomidine is better than midazolam for premedication in children as it produces better sedation, parental separation and satisfactory ease of induction by successful mask acceptance.

Keywords: Dexmedetomidine, Intranasal, Midazolam, Mucosal Atomization Device, Pediatric

Introduction

The preoperative period is a stressful time for children<sup>2</sup>. It has been observed that preoperative anxiety of parental separation predisposes children to sleep disturbances and behavioural changes postoperatively. Zeev N kain et al., suggested that the prevalence of preoperative anxiety was high and reported to range from 40 - 60% among young children before anesthesia induction and surgery<sup>3</sup>. This is a concern for anaesthesiologists.

An ideal premedicant relieves anxiety, makes the children calm, reduces their fear, makes induction smooth, rapid recovery, provides good patient acceptance and parental separation<sup>4</sup>.

In pediatric patients benzodiazepines are commonly used for premedication as they provide sedation, muscle relaxation, anxiolysis, hypnosis, amnesia and anticonvulsant properties.

Midazolam is water soluble short acting gammaamino butyric acid receptor inhibitor. It has faster onset of action. Thus gained popularity as premedication in children<sup>5</sup>.

Dexmedetomidine has been explored extensively in the pediatric population. It is highly selective and specific agonist for  $\alpha_2$  adrenoceptor exhibits sedative, hypnotic, analgesic, anxiolytics and sympatholytic effect. It has minimal effect on respiratory drive. These properties render dexmedetomidine suitable for analgesia and sedation during preoperative period<sup>6</sup>.

Children dislike forcible administration of injection which will lead to struggling and psychological impact. Intravenous and intramuscular routes cause more anxiety. Better acceptability of intranasal route is considered for premedication in children as it is non invasive route and its ease of administration. Large and well vascularized nasopharyngeal mucosal surface provided rapid absorption, early onset via intranasal route with high bioavailability. It had also the advantage of well tolerability, did not require children's cooperation as would be in case for drug swallowing or sublingual retention and did not have pungency or an unpleasant taste<sup>8</sup>. Atomization of drug intranasally by Mucosal Atomization Device (MAD) produced fine particles (30-100 micron in diameter) which was associated with less discomfort during administration and increase drug absorption<sup>9</sup>. So, we carried out a randomised study of midazolam

versus dexmedetomidine for premedication in children via intranasal Mucosal Atomization Device.

### **Material And Methods:**

This randomised study was carried out in the department of Anaesthesiology at a tertiary health care Centre after obtaining approval from the institutional ethical committee. A total of 60 children of either sex of 2-6 years belonging to ASA I or II, scheduled for elective surgeries under general anesthesia were included in the study. Upper respiratory tract infections, any nasal pathology, any known allergy or sensitivity or any other form of reaction to benzodiazepines and  $\alpha_2$  adrenoceptor agonists, children with mental retardation, on anticonvulsant therapy or other sedative medications were excluded from the study.

Detailed preanaesthetic history was taken a day prior to surgery. General examination, physical and systemic examination and thorough airway assessment were carried out. Children's weight were noted. All routine investigations were done.

We explained the procedure, the device, drugs and details about their administration, their probable side effects to parents. Written parental informed consent was taken in their native language. Children were kept nil by mouth for solids about 6 hours and clear fluids were permitted upto 2 hours prior to the surgery. The children were kept in quiet, undisturbed area along with the parents.

Primary parameters included were:

Acceptance of the drug was assessed by using the **Drug Acceptance Score** (Parnis S.J. et al)<sup>10</sup>

- 1. Rejected entirely.
- 2. Accepted with grimace or complaint.
- 3. Accepted readily

Sedation was assessed at 30 minutes after the administration of study drug by **Four Point Sedation Score (Filos et al)**<sup>11</sup>

- 1. Awake and alert,
- 2. Awake but drowsy, responding to verbal stimuli,
- 3. Drowsy but responding physical stimulus,
- 4. Unresponsive, not responding to physical stimulus.

The behaviour at the time of separation from parents was assessed when the child was separated from parents to shift to operating room using the **Parental** Separation Score (Pandit UA et al)<sup>12</sup>

- 1. Excellent, happily separated,
- 2. Good, separated without crying,
- 3. Fair, separated with crying,
- 4. Poor, need for restraint.

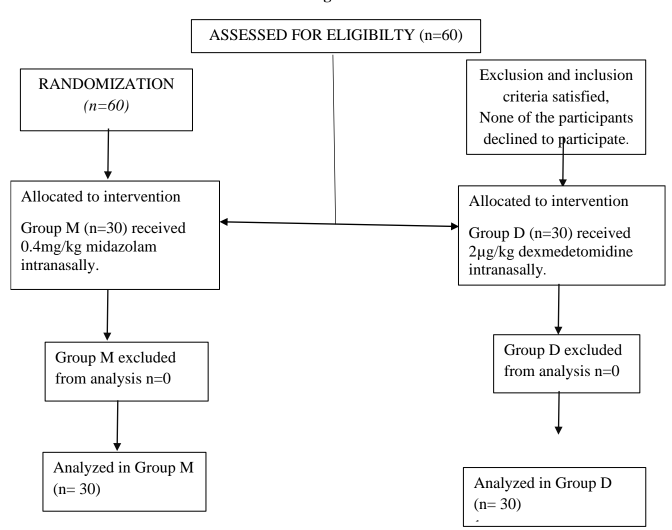
Acceptance of face mask was graded on <u>Four Point</u> <u>Score (Mitchell V)</u><sup>13</sup>

- 1. Poor afraid, combative, crying,
- 2. Fair moderate fear of mask, not easily calmed,

- 3. Good slight fear of mask, easily calmed,
- 4. Excellent unafraid, cooperative, accepts mask easily.

Secondary parameters included baseline pulse rate,  $SpO_2$  and monitored every 15 minutes till the end of surgery. Sample size had been estimated with the help of statistical software nMaster 2.0., sample size came out as 30 patients in each group. They were divided equally into 2 groups. Children in Group M received a dose of 0.4 mg/kg midazolam (upto a maximum of 10mg) and children in Group D received dexmedetomidine in a dose of  $2\mu g/kg$  (upto a maximum of  $50 \mu g$ ).

Figure 1



The half of the calculated dose of the drug was administered in each nostril 30 minutes before surgery in a recumbent position using mucosal atomization device by an experienced anaesthesiologist. Children were shifted to operation theatre and were premedicated with injection

glycopyrrolate 0.004mg/kg intravenous and injection ondansetron 0.1mg/kg intravenous. They were preoxygenated with face mask with 100% oxygen for 3 minutes. Anaesthesia was induced by a standard technique of intravenous induction and maintained on  $N_2O$ . sevoflurane atracurium.  $O_2$ and Intraoperatively children were monitored for pulse rate, SpO<sub>2</sub> every 15 minutes till end of surgery. At the end of surgery neuromuscular blockade was reversed with inj.neostigmine (0.05mg/kg) and inj. glycopyrrolate (0.008mg/kg). Trachea was extubated after fulfilling the recovery criteria and children were shifted to recovery room. Postoperatively all children were watched for pulse rate, SpO<sub>2</sub>, nausea, vomiting, rigor, bradycardia every hourly up to 4 hours and at 6 and 8 hours.

Bradycardia was defined as pulse rate < 60/min and treated with IV atropine sulfate 0.6mg.

### **Statistical Analysis:**

Data was collected, tabulated. Numerical variables were presented as mean and standard deviation (SD) while categorical variables were presented as frequency and percentage. As regard numerical variables, unpaired student — t test was used whenever appropriate between-group comparisons; while for categorical variables, chi—square test was used. A difference with significant level (p<0.05) was considered statistically significant.

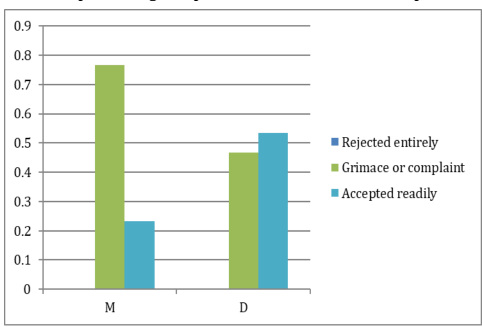
### **Observation And Results:**

Demographically both the groups were comparable on the basis of age, gender, weight and ASA grading.

		8 1		8 1	
Drug Acceptance	Group M		Group D		р
Score	No of pts.	%	No of pts.	%	value
1- Rejected entirely	0	0.00%	0	0.00%	
2- Grimace or complaint	23	76.67%	14	46.67 %	0.033
3- Accepted readily	7	23.33%	16	53.33%	] '
Total	30	100.00%	30	100.00%	

Table 1 - Drug Acceptance Score between the groups:-

Based on the above results it was found that the **Drug Acceptance Score** in Group D (dexmedetomidine  $2\mu g/kg$ ) was statistically significantly good as compared to Group M (midazolam 0.4mg/kg) (p=0.0337). None of the children in any group rejected the drug.

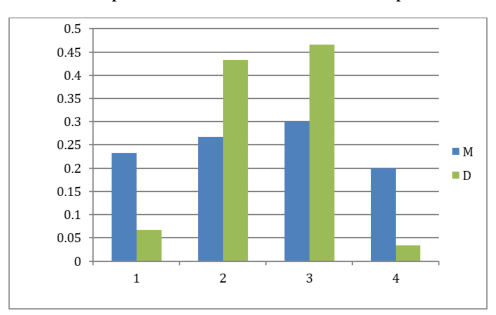


**Graph 1 - Drug Acceptance Score Between The Groups:** 

Table 2 - Sedation Scores Between The Groups:-

	Group M		Group D		
Four Point Sedation Score		%	No of pts.	%	p value
1- Awake and alert	7	23.33%	2	6.67%	
2- Awake but drowsy,responding to verbal stimuli	8	26.67%	13	43.33%	
3- Drowsy but responding physical stimulus	9	30.00%	14	46.67%	0.0347
4- Unresponsive	6	20.00%	1	3.33%	
Total	30	100.00	30	100.00	

Based on the various studies we had compared Sedation Score at 30 minutes after administration of the study drug. The Sedation Score was statistically significant in Group D when compared to Group M (p=0.0347).

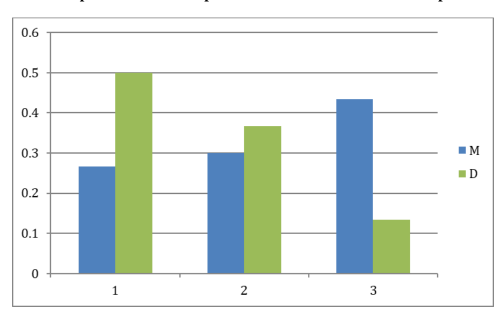


**Graph 2 - Sedation Scores Between The Groups** 

Table 3 – Parental Separation Score Between The Groups:-

Parental Seperation Score	Group M		Group D		p value
	No of pts.	%	No of pts.	%	r ·······
1- Excellent, happily separated	8	26.67%	15	50.00%	
2- Good, separated without crying	9	30.00%	11	36.67%	0.0288
3- Fair, separated with crying	13	43.33%	4	13.33%	0.0288
4- Poor, need for restraint	0	0.00%	0	0.00%	
Total	30	100.00%	30	100.00%	

In the present study overall **Parental Separation Score** was statistically significantly good to excellent in dexmedetomidine group as compared to midazolam group (p = 0.0288). None of the children in any group were needed to be restrained.

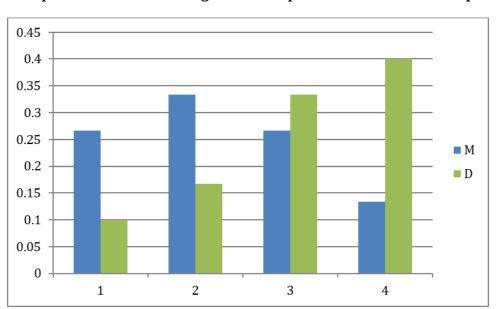


**Graph 3 – Parental Separation Score Between The Groups** 

Table 4 - Behaviour During Mask Acceptance Between The Groups (FOUR POINT SCORE):-

Acceptance of	Grou	рМ	Grou	p value	
Face Mask	No of pts.	%	No of pts.	%	_ p varae
1- Poor	8	26.67%	3	10.00%	
2- Fair	10	33.33%	5	16.67%	
3- Good	8	26.67%	10	33.33%	0.0428
4- Excellent	4	13.33%	12	40.00%	
Total	30	100.00%	30	100.00%	

In the operation theatre the acceptance of mask (**Four Point Score**) was compared as it has an impact on induction. The overall **Mask Acceptance Score** was statistically significantly good to excellent in Group D as compared to Group M (p value = 0.0428).



**Graph 4 - Behaviour During Mask Acceptance Between The Groups** 

Table 5

Publication Author	Type of study	Study population number and age.	Drugs and Method	Results
Malineni N. et al (2017) <sup>1</sup>	Randomise d controlled trial, double blind study	60 children. 1-10 years.	<ul> <li>Midazolam 0.2mg/kg.</li> <li>Dexmedetomidine 1µg/kg.</li> <li>Administered with 1ml tuberculin syringe intranasally to assess parental seperation anxiety and acceptance of anaesthesia mask.</li> </ul>	Parental seperation and mask acceptance at time of induction was better in dexmedetomidine when compared to midazolam.
Kumar A. et al (2017) <sup>18</sup>	Prospective , randomised , double	60 children 2-12 years.	- Oral midazolam 0.5mg/kg - Intranasal	Sedation scores were superior with dexmedetomidine group at separation and

	Γ	T		
	blind study		dexmedetomidine1µg/kg	induction.
			- 0.2ml drug dripped into both nostrils using 1 ml	
			syringe to assess sedation	
			scores at seperation from	
			parents and at induction of anaesthesia.	
			anaestnesia.	
Xie z. et al	Randomise	106 children	Dexmedetomidine 2µg//kg	Dexmedetomidine by
	d study	100 cilidren	in 20µl//kg of Normal	MAD offered better
$(2017)^{19}$		2-5 years.	saline is given as drops via	sedation effects to
			syringe or spray using	reduce responses to
			MAD to assess response to	venous cannulation than
			venous cannulation by FLACC scores (faces,	by drops without any significant
			legs, activity, cry and	complications.
			consolability) after 30	-
			minutes of administration.	
Gupta A. et al (2017) <sup>20</sup>	Prospective	60 children	- Midazolam 0.2mg/kg.	Dexmedetomidine resulted in more
(2017)	, randomised	1-8 years.	- Dexmedetomidine	successful parental
	, double		1μg/kg	separation and yielded a
	blind study		- Administered	higher sedation level
			intranasally using 1ml	than midazolam.
			syringe to assess time of	
			onset, level, sedation quality upon separation	
			from parents.	
			r	

Naik Shilpa S. et al (2018) <sup>21</sup>	Longitudin al study	30 children 1-5 years.	- Midazolam spray intranasally 0.5mg/kg.  - Dexmedtomidine instillation intranasally 1μg/kg.  - To assess compliance to intravenous cannulation, separation from parents and induction score, sedation score, postoperative recovery score.	Dexmedetomidine spray gave gives better sedative condition, response to i.v. cannulation, separation and induction as compared to midazolam.
Medhat MM. et al (2018) <sup>14</sup>	Prospective , randomised , double blind study	60 children 3-6 years.	- Midazolam 0.2mg/kg  - Dexmedetomidine 1μg/kg  - By drop instillation intranasally using 2ml syringe to assess sedation score, anxiety score and child- parent separation score.	Dexmedetomidine attained satisfactory and significant sedation and lower anxiety level with better parental separation than intranasal midazolam.

Diwan G.et al	Prospective	60 children	- Midazolam 0.1mg/kg	Dexmedtomidine was
$(2020)^{22}$	randomised , double blind study	2-12 years.	- Dexmedetomidine 1μg/kg  - Administered intranasally using 1ml syringe comes to assess sedation score, anxiety score and child- parent separation score.	associated with lower sedation levels, anxiety levels and easier child parent seperation when shifting to operating room when compared to midazolam group.

In both the groups intraoperative and postoperative heart rates were found to be statistically insignificant (p>0.05). No medical intervention was needed in either group.

Nasopharyngeal irritation was not seen in any patients in our study.

No significant change was observed in SpO2 between both the groups intraoperatively as well as postoperatively. None of the patients in both groups had SpO2 < 98% at any point of time during study period.

In our study none of the children in both groups had complications such as bradycardia, hypertension and hypotension both during intraoperative and postoperative period.

### **Discussion:**

Pediatric patients undergoing surgical procedures can experience significant anxiety and distress during perioperative period. They are usually uncooperative, anxious or physically resistant particularly during the times of parental separation, mask application and venipuncture<sup>4</sup>. Various interventions were used to allay the anxiety of a child during perioperative preparation. Sedation in preoperative room remained one of the widely used method and helped to reduce

anxiety, minimized the emotional trauma and facilitated a smooth induction of anaesthesia<sup>14</sup>.

The intranasal application of pre anaesthetic drugs is a preferred route of administration and is an effective way to administer sedatives to children. It doesn't require cooperation and it is convenient, noninvasive, well tolerated. Child would not be having an unpleasant taste or pungency.

The Mucosal Atomization Device is safe simple metered dose delivery system and painless way to deliver medications as it is needle free. Atomization can be done in any position. The soft conical plug on the tip forms a seal with the nostril, preventing expulsion of drug. Atomised nasal medications are the optimal size for rapid absorption across mucosal membranes into the blood stream, avoiding first pass metabolism<sup>15</sup>.

We compared the intranasal administration of midazolam (0.4mg/kg) and dexmedetomidine  $(2\mu g/kg)$  and as premedication in pediatric age group undergoing elective surgeries under general anaesthesia using MAD.

Midazolam which is a water soluble benzodiazepine has emerged as a widely used pre medication due to its fast onset of action and short elimination half life<sup>3</sup>. It binds to GABA<sub>A</sub> receptor triggering chloride

channel and hyperpolarization of cells thus causing resistance to excitation of neuron, hence producing sedation.

Intranasal administration of midazolam was better tolerated than oral and has the advantage of no first-pass effect with rapid absorption directly into the systemic circulation and a bioavailability of 55-83% <sup>16</sup>.

Dexmedetomidine is a  $\alpha_2$  agonist which produces cooperative sedation and no respiratory depression. It produces its hypnotic and sedative effects by neuronal hyperpolarisation via activation of  $\alpha_{2A}$  adrenoceptor on the predominant noradrenergic nucleus in the brain, locus coeruleus.

Intranasal dexmedetomidine administration had a high bioavailability of 35-93% (65%)<sup>17</sup>

### **Conclusion:**

We conclude that administration of intranasal dexmedetomidine  $2\mu g/kg$  is better than intranasal midazolam 0.4mg/kg for premedication in pediatric age group between ages of 2 and 6 years via Mucosal Atomization Device as it produces satisfactory ease of drug acceptance, sedation, parental separation and induction by successful mask acceptance without causing any complications.

#### **References:**

- Malineni N, Patil MC. "Comparative evaluation of intranasal dexmedetomidine and intranasal midazolam for premedication in children undergoing anesthesia": A 1-year double-blind randomized controlled trial. Indian J Health Sci Biomed Res 2017:10:155-9.
- 2. Kain ZN, Mayes LC, O'Connor TZ, Cicchetti DV. Preoperative anxiety in children: predictors and outcomes. Arch Pediatr Adolesc Med 1996; 150: 1238–45.

- 3. Kain ZN, Mayes LC, Caldwell-Andrews AA, Karas DE, McClain BC. Preoperative anxiety, postoperative pain, and behavioral recovery in young children undergoing surgery. Pediatrics. 2006 Aug; 118(2):651-8. doi: 10.1542/peds.2005-2920. PMID: 16882820.
- 4. Creedon LR, Dock M. Pharmacological management of patient behavior. In: McDonald RE, Avery DR, editors. Dentistry for the Children and Adolescent. 8th ed. St. Louis: CV Mosby; 2004. p. 285-311.
- 5. Tschirch FT,Gopfert K,Frohlich JM,Bruner G,Weishaupt D.Low-dose intranasal versus oral midazolam for routine body MRI of claustrophobic patients. Eur Radiol.2007;17:1403-10.[PubMed][Google Scholar].
- 6. Talon MD, Woodson LC, Sherwood ER, Aarsland A,McRae L, Benham T, et al.Intranasal dexmedetomidine premedication is comparable with midazolam in burn children undergoing reconstructive surgery.J Burn Care Res.2009;30:599- 605.[PubMed][Google Scholar].
- 7. Wang J,Bu G.Influence ofss intranasal medication on the structure of nasal mucosa.Chin Med J(Engl)2002;115:617-9.[PubMed][Google Scholar].
- 8. Yuen VM, Hui TW,Irwin MG,Yao TJ,Wong GL, Yuen MK, et al.Optimal timing for the administration of intranasal dexmedetomidine for premedication in children.Anaesthesia.2010;65:922-9.[PubMed][Google Scholar].
- 9. Wolfe TR, Braude DA. Intranasal medication delivery for children: a brief review and update. Pediatrics 2010;126:532-7. Crossers PubMed Web of Science R Google Scholar.

- 10. Parnis SJ, Foate JA, van der Walt JH, Short T, Crowe CE. Oral midazolam is an effective premedication for children having day-stay anaesthesia. Anaesth Intensive Care. 1992 Feb;20(1):9-14. doi: 10.1177/0310057X9202000102. PMID: 1609951.
- 11. Filos KS, Goudas LC, Patroni O, Polyzou V. Hemodynamic and analgesic profile after intrathecal clonidine in humans. A dose-response study.

  Anesthesiology.1994;81:591-601.[PubMed][Google Scholar].
- 12. Pandit UA, Collier PJ, Malviya S, VoepelLewis T, Wagner D, Siewert MJ. Oral transmucosal midazolam premedication for preschool children. Can J Anaesth 2001;48:191-5.
- Mitchell V, Grange C, Black A, Train J. A comparison of midazolam with trimeprazine as an oral premedicant for children. Anaesthesia 1997;52:416
- Medhat MM and Gamal Zakaria El-Morsy. Anaesth Essays Res. 2018 Jan- Mar; 12(1): 170-175. Doi:10.4103/aer.AER 119 17.PMID: 29628576.
- 15. Corrigan M, Wilson S, Hampton J. Safety and efficacy of intranasally administered medications in the emergency department and prehospital settings. Am J Health-Syst Pharm. 2015; 72: 1544-1554.
- 16. Björkman S, Rigemar G, Idvall J. Pharmacokinetics of midazolam given as an intranasal spray to adult surgical patients. Br J Anaesth. 1997;79:575–80. PubMed.
- 17. Yildirim SV, Guc BU, Bozdogan N, Tokel K. Oral versus intranasal midazolam premedication

- for infants during echocardiographic study. Adv Ther. 2006;23:719–24. PubMed.
- 18. Kumar A, Sinha C, Kumar A, Kumari P. The effect of intravenous dexmedetomidine compared to propofol on patients hemodynamics as a sedative in brachial plexus block: A comparative study. Anesth Essays Res 2017;11:201-5.
- 19. Xie Z, Shen W, Lin J, Xiao L, Liao M, Gan X. Sedation effects of intranasal dexmedetomidine delivered as sprays versus drops on pediatric response to venous cannulation. Am J Emerg Med. 2017 Aug;35(8):1126-1130. doi: 10.1016/j.ajem.2017.03.021. Epub 2017 Mar 18. PMID: 28347608.
- 20. Gupta A, Dalvi NP, Tendolkar BA. Comparison between intranasal dexmedetomidine intranasal midazolam as premedication for brain magnetic resonance imaging in pediatric patients: A prospective randomized double blind trial. J Anaesthesiol Clin Pharmacol. 2017 Apr-Jun:33(2):236-240. doi: 10.4103/joacp.JOACP 204 16. PMID: 28781452; PMCID: PMC5520599.
- 21. Naik Shilpa S, Shrama Parul. To assess the Compliance and Sedation Score of Intranasal Midazolam and dexmedetomidine Premedication among Children. Indian J Anesth Analg. 2018;5(12):2062-66.
- 22. Diwan G, Bharti AK, Rastogi K, Gupta PK. Comparison of intranasal dexmedetomidine and midazolam as premedication in pediatric surgical patients: A prospective, randomized double-blind study. Anesth Essays Res 2020;14:384-9.