COMPARATIVE STUDY TO EVALUATE THE EFFECTS OF INTRANASAL DEXMEDETOMIDINE VERSUS MIDAZOLAM AS A PREMEDICATION AGENT IN CHILDREN UNDERGOING ELECTIVE SURGERY

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ABSTRACT

BACKGROUND
Intranasal route of drug administration is a practical option as a non-invasive alternative to intravenous route especially for children. We conducted a study to compare the effects of Dexmedetomidine and Midazolam administered as premedication agents through intranasal route. The primary outcome measures were assessing the sedation status upon parental separation and behaviour level during induction. The secondary outcome measures were assessing the acceptance of intravenous cannulation and acceptance of facemask during induction.

MATERIALS AND METHODS
Hundred children aged between 5–12 years of both sexes belonging to ASA grade I and II and weighing <30 kg were enrolled in this prospective randomised double blinded study. The children were divided into two groups of 50 each. 60 minutes before induction, Group IN-D (n = 50) received Intranasal Dexmedetomidine 1 µg/ kg and Group IN-M (n = 50) received Intranasal Midazolam 0.2 mg/kg.

RESULTS
Sedation status at parental separation produced by dexmedetomidine is statistically significant when compared to midazolam. Both the drugs produced similar behaviour level inside the operating room, acceptance of IV cannulation and acceptance of facemask during induction.

CONCLUSION
Intranasal dexmedetomidine can be used as a preanaesthetic medication in children undergoing elective surgeries under general anaesthesia.

KEYWORDS
Anaesthesia, Premedication, Intranasal, Dexmedetomidine, Midazolam.


BACKGROUND
Preoperative anxiety is a subjective feeling of worry, tension, apprehension and nervousness. In children, it is attributed to separation from parents.1 It largely affects the smoothness of induction, emergence from anaesthesia and also the psychological and emotional state of children in the remote future.2 A smooth transition from an awake state to surgical anaesthesia in children becomes a challenge for all anaesthesiologists. Eliminating or minimising stress provides a smooth atraumatic induction of anaesthesia. Preoperative preparation programs, parental presence during induction and premedication with drugs are methods to allay the anxiety of the children in the perioperative period.3

Among the various routes of premedication, intranasal route has come into practice from early nineties.4,5 It is relatively a quick, painless with a high bioavailability. Most commonly used drug for premedication is Midazolam. Dexmedetomidine, an α2 agonist, is now becoming popular because of its excellent sedative, anxiolytic, sympatholytic and analgesic properties.6,7,8

This study was conducted to compare the effects of Dexmedetomidine and Midazolam administered intranasally for premedication in children undergoing elective surgery under general anaesthesia.

MATERIALS AND METHODS
This study was conducted in Thanjavur Medical College Hospital from August 2011 to October 2013. 43 was the estimated sample size in each group. According to a study by Meenakshi Sundaram AL et al.,9 at induction of anaesthesia, 24.7% and 67.4% of the children from groups M and D...
respectively were satisfactorily sedated with an allocation ratio of 1, Alpha = 0.01 and Beta = 0.10. Finally, a total of 100 children scheduled for elective surgeries under General Anaesthesia were included, 50 in each group. Children meeting the following selection criteria were included in the study. Block randomisation technique was used to allocate the children to midazolam and dexmedetomidine group. Blocks of varying sizes ranging from 2 - 6 used for block randomisation. Approval from Institutional Ethical Committee was obtained.

Inclusion Criteria
1. Children of both sexes aged 5 – 12 Yrs.,
2. Belonging to ASA I and II,
3. Weighing <30 kg,
4. Undergoing elective surgery under GA.

Exclusion Criteria
1. Known allergy or Hypersensitivity to Dexmedetomidine or Midazolam,
2. Organ dysfunction,
3. Cardiac arrhythmia,
4. Congenital heart disease,
5. Mental retardation,
6. Patient refusal,
7. H/o snoring or sleep apnoea,

The Children were Divided into Two Groups-
Group IN-D (n = 50) - Intranasal Dexmedetomidine.
Group IN-M (n = 50) - Intranasal Midazolam.

Children were kept nil per oral for 8 hours. Informed written consent was obtained from the parents. Children were randomly assigned to one of two groups. Baseline heart rate, systolic blood pressure and SPO2 were recorded.

Group IN-D received 1 µg/kg of intranasal dexmedetomidine and Group IN-M received 0.2 mg/kg of intranasal midazolam 60 minutes before induction.

The study drugs were prepared by an independent person using the intravenous preparations of dexmedetomidine (100 µg/kg) and midazolam (5 mg/mL). Intranasal medication was prepared in a 2-ml syringe and 0.9% saline was added to make a final volume of 1.5 mL. The medication was administered in preoperative room in the presence of one parent (Preferably mother). The drug was dripped into the nostrils with the child in the recumbent position. Patients and the observer were blinded to the drug being administered.

Continuous heart rate and SPO2 monitoring done. Desaturation was managed with O2 supplementation through nasal mask. After 60 minutes the child was shifted to the operating table.

Sedation status was assessed during parental separation with a 6 - point sedation scale Modified from the Observer Assessment of Alertness and Sedation scale.
6 - Appears alert and awake, responds readily to name.
5 - Appears asleep, responds readily to name spoken in normal tone.
4 - Lethargic response to name spoken in normal tone.
3 - Respond only after name called loudly or repeatedly.
2 - Responds only to mild prodding or shaking.
1 - Does not respond to mild prodding or shaking.

Sedation score from 1 - 4 was considered satisfactory while score from 5 – 6 was considered unsatisfactory.

Behaviour score during induction was assessed on the operating table by a 4 - point scale-
4 - Crying or resisting.
3 - Anxious and could not be reassured.
2 - Anxious but could be reassured.
1 - Calm and co-operative.

Behaviour score of 1 or 2 was considered satisfactory while 3 or 4 was considered unsatisfactory.

The acceptance of IV cannulation in the forearm of the child was graded as-
Grade 1 - Calm, co-operative, asleep - Good.
Grade 2 - Co-operative with reassurance - Moderate.
Grade 3 - Combative, crying - Poor.

Glycopyrrolate 10 µg/kg, ondansetron 0.15 mg/kg were given IV preoxygenation done with 100% O2 for 3 minutes with a scented facemask. Acceptance of facemask by the child was graded with the same grading as that of the acceptance of IV cannulation.

With standard monitoring, General Anaesthesia was administered. Recovery was good and the children were shifted to postoperative ward.

Statistical Analysis
The collected data were analysed by using GraphPad Instat 3.06 software according to variables for chi square test and student's t-test. The results are obtained in the form of range, mean and standard deviation. The probability value 'p' of less than 0.05 was considered statistically significant.

RESULTS
Patient's demographic data including age, sex and weight between two groups were comparable (table 1). The age distribution was in the range of 5 - 12 years in both the groups. The mean age of group IN-D is 9.04 and group IN-M is 8.80. The mean weight of group IN - D was 23.36 kg and that of group IN-M was 23 kg. The p value was 0.692 (>0.05) was statistically not significant.

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Data</th>
<th>Group IN - D</th>
<th>Group IN - M</th>
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<td>1.</td>
<td>Age (Mean)</td>
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<td>8.80</td>
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<tr>
<td>2.</td>
<td>Weight in Kgs. (Mean)</td>
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<td>3.</td>
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<tr>
<td></td>
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Table 1. Demographic Data

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<th>Parameter</th>
<th>Age in years</th>
<th>Weight in kilograms</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Group IN-D (n = 50)</td>
<td>Group IN-M (n = 50)</td>
</tr>
<tr>
<td>Range</td>
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<td>5-12</td>
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<tr>
<td>Mean</td>
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</tr>
<tr>
<td>S.D.</td>
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<td>1.807</td>
</tr>
<tr>
<td>'p' value</td>
<td>0.4227 (&gt;0.05)</td>
<td>0.692 (&gt;0.05)</td>
</tr>
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</table>

Table 2. Age and Weight Distribution
The mean sedation score in group IN-D was 3.24 and in group IN-M was 4.32. Sedation score at parental separation was satisfactory in 88% (44 children) of the group IN-D and in 60% (30 children) of the group IN-M. It was found to be unsatisfactory in 12% (6 children) of the group IN-D and in 40% (20 children) of the IN-M group. The ‘p’ value was 0.030 (<0.05) which was statistically significant.

Acceptance of facemask was good in 68% of the children in group IN-D and in 60% of the children in group IN-M. Only two children (4%) in group IN-D had a poor acceptance of facemask (‘p’ = 0.401).

The table 3 shows the distribution of acceptance of IV cannulation between two groups. The grading was good in 36%, moderate in 56% and poor in 8% of the children in both the groups.
Continuous heart rate and SpO2 monitored, none of the children in either of the groups attained significant bradycardia and desaturation (<95%). No subjective adverse effects were observed in group IN-D. Three children complained of nasal irritation and child had paradoxical excitation in group IN-M following drug administration. These parameters were not statistically analysed as our study was not designed to investigate these parameters.

**DISCUSSION**

Selecting appropriate premedication in safe yet effective doses is a primary responsibility of an anaesthesiologist. Facilitating an anxiety free separation and a smooth induction are the primary goals of premedicant drug administration. Analgesia, amnesia, and prevention of physiologic stress are other beneficial effects of pharmacological intervention. The main drawback of premedication particularly in children is that the process of administering the medication may actually increase anxiety. A child’s major fear associated with hospitalisation is fear from needles and injections. Thus, a non-invasive route is generally preferred.

Various routes for premedication in children have been formulated keeping in mind the specific problems of the paediatric age group. Intranasal route is a rapid and effective way of administering premedication in children. Among the non-invasive routes, it has gained importance in recent decades. It provides a direct route into the systemic circulation for drugs that easily cross mucous membranes. Importantly the first bypass metabolism is circumvented and the bioavailability is better than other routes. For many intranasal medications, the rates of absorption and plasma concentrations are comparable to intravenous administration and are typically better than subcutaneous or intramuscular routes. Oral route is poorly accepted and rectal route is less ideal especially for older children.

Alpha - 2 agonists are assuming greater importance as anaesthetic adjuvants and analgesics.10,11 Dexmedetomidine is a highly selective, specific and potent alpha - 2 adrenoceptor. It was introduced in United States and had been in clinical practice since 1999. Sedative - hypnotic effects are produced by the action on alpha - 2 receptors in the locus coeruleus. Analgesic effect is produced by its action on alpha - 2 receptors within the locus coeruleus and within the spinal cord. Despite effective sedation there is limited respiratory depression. Following intranasal administration, 1 µg/kg has an onset time of 45 minutes with a peak effect at 60 - 105 minutes in healthy adults. Bioavailability of intranasal dexmedetomidine is 65% (35 - 93%).12 Whereas the sedation, anterograde amnesia and anticonvulsant properties of midazolam are mediated via alpha 1 GABA receptors. These receptors are found in highest densities in the olfactory bulb, cerebral cortex, cerebellum, hippocampus, substantia nigra and inferior colliculus.

Among the large percentage of paediatric patients, anaesthesia induction was known to be the anxiety provoking part. Awake intravenous cannulation and parental separation were more vulnerable points. So this study was designed to compare the efficacy of intranasal dexmedetomidine and midazolam with respect to sedation status at parental separation, behaviour level inside the operating room, acceptance of IV cannulation and acceptance of facemask during induction.

Several studies have used intranasal midazolam and the possibility of neurotoxicity occurred only after chronic administration through intrathecal route.13 Timiliolra et al12 showed intranasal dexmedetomidine does not produce local effects on nasal mucosa like mucosal irritation, ulceration, inflammation and bleeding. Pradipabakata et al14 in their study concluded that intranasal midazolam in a dose of 0.2 mg/kg was as effective as 0.3 mg/kg in producing anaesthesia and sedation. Yuen et al15 observed that 1 µg/kg dexmedetomidine produced significant and satisfactory sedation at parental separation and at induction. Aynur et al16 compared the efficacy of 1 µg/kg of dexmedetomidine and 0.2 mg/kg of midazolam administered intranasally as premedication agents in children between 2-9 years.

The sedation produced by dexmedetomidine significantly differs when compared with other drugs that act through GABA systems.15 Among the 50 children, 22 children (88%) in group IN-D and 15 children (60%) in group IN-M attained a significant and satisfactory sedation status at parental separation at 60 minutes (p = 0.030). While Aynur akin et al16 showed that 79.9% in group D and 95.5% in group M attained satisfactory sedation (45 minutes). The peak effect of intranasal midazolam is 10-15 minutes and duration of action is 30-60 minutes. In this study, the sedation status was assessed at 60 minutes, so only 60% in group IN-M was found with satisfactory sedation. The mean sedation score in group IN-D was 3.24 ± 0.959 and in group IN-M was 4.32 ± 0.740 (p = 0.001).

The behaviour level of the children was assessed with a four point scale. 80% of children in group IN-D and 92% in group IN-M attained satisfactory behaviour score (score 1 or 2). None of the children in either groups had worst and unsatisfactory behaviour (score 4). 3 children (12%) in group IN-D and 2 (8%) in group IN-M had score 3 (Anxious, not reassured). Yuen et al15 in their study also showed that children in groups M, D-0.5 and D-1 premedicated with midazolam (0.5 mg/kg) and dexmedetomidine (0.5 and 1 µg/kg) had comparable behaviour level. There was a tendency of the children in groups D-0.5 and D-1 having unsatisfactory behaviour after shifting to operating room. But still there were no statistically significant differences when compared to group M.

Children usually have an aversion for needles. If acceptance of the IV cannulation is good enough, it indirectly counterchecks the sedation status and behaviour level. The
children in both groups IN-D and IN-M had similar acceptance of IV cannulation (p = 1.000).

The acceptance of facemask was good (score 1) in 68% of the children in group IN-D and in 60% of the children in group IN-M. It was found to be moderate (score 2) in 28% of the children in group IN-D and in 40% of the children in group IN-M. 2 children (4%) in group IN-D had a poor (score 3) acceptance of facemask (p = 0.401).

Limitations of the Study
This study was conducted entirely in the preoperative period, the intraoperative and the postoperative effects were not evaluated.

In summary, we found that intranasal administration of dexmedetomidine 1 µg/kg for premedication in children undergoing elective surgery provides significant and satisfactory sedation (p’ = 0.030). However, intranasal midazolam was as effective as dexmedetomidine in providing satisfactory behaviour level inside the operating room. Both the intranasal premedicants were equally effective in providing satisfactory conditions for acceptance of IV cannulation and acceptance of facemask during induction.

CONCLUSION
Intranasal premedication allows an effective and predictable sedation in children. Both midazolam and dexmedetomidine produce a good level of sedation, behaviour and acceptable levels for IV cannulation and face mask acceptance during induction. But the quality of sedation is significantly better in dexmedetomidine group.

REFERENCES