Original Article

Premedication of Oral Clonidine 2mcg/kg and 4mcg/kg for Analgesic and Pressure Response During First Twenty-four Hours After Upper Abdominal Surgery: A Comparative Study


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Context background:
Traditionally postoperative period analgesia was maintained by opioids and which frequently increase the hospital stay time and cost by delayed bowel movement and others side effects. on the other hand upper abdominal surgery is very much painful, usually Opiods are used as good analgesic, but have some adverse effect and addiction effect; anaesthesiologists want to reduce its requirements. In upper abdominal surgery adequate analgesia, stable haemodynamic, early bowel movement, free from nausea and vomiting is wanted, as a part of multimodal analgesic approach, premedication by clonidine is very important for it analgesic, angiolytic and sedative properties. Alpha two (α-2) adrenoreceptor agonist, Clonidine exerts central sympatholytic effect for 8 to 10 hours as its half life is 9-12 h. So that premedication with oral clonidine causes reduction of anxiety, reduction of perioperative analgesic drugs and also reduction of anaesthetic doses. In addition, clonidine increases cardiac baroreceptor reflex sensitivity to increase in systolic blood pressure, and thus stabilizes blood pressure. Clonidine is rapidly and almost completely absorbed after oral administration with a time to maximum plasma concentration of between 1.5 and 2 hr and elimination half-life of 8 to 12 hr. But clonidine produces analgesia in a dose dependent manner, achieving complete pain relief for up to 5 hours without sensory or motor block at large doses (oral 7000 to 900 mcg) however large doses were associated with disadvantage including hypotension, bradycardia and transient sedation. It also reported that clonidine 150mcg intravenous (I/V) produce a similar analgesic effect to morphine 5mg in patient after orthopedic surgery. Because of its dose, route, and surgical variation it is very much important to specify the dose for upper abdominal surgery.

The primary aim of this study to compare the effects of clonidine premedication at different doses (2 & 4mcg/kg) on postoperative analgesia and hemodynamic status in upper abdominal surgery with large incision area. (like- hepatobiliary surgery, gastrectomy, esophagectomy, hepaticectomy, and whipples operations).

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Participants: 60(sixty) v patients was selected from the population by non probability sampling technique. All patients with ASA class-I, & class-II were selected after they are admitted in Dhaka medical college hospital, surgery department for elective upper abdominal surgery (like-gastrectomy, pancreatectomy, hepaticotomy, open cholecystectomy, spleenectomy, hepatobiliary surgery, bariatric surgery). Before selection all patients' history, investigation and examination were done according to the inclusion and exclusion criteria. Patients included age 18 to 60 years, undergoing upper abdominal surgery with ASA physical status I, II, and those who gave informed consent were eligible for this study. The potential participants were approach by researcher one week before the admission in hospital for operation. Exclusion criteria hypersensitivity to clonidine or any of the drugs in operation and history of uncontrolled hypertension, cardiovascular co-morbidities, known un controlled diabetic with diabetic neuropathy, alcohol abuse, opium addict or any drug that modifies pain perception, Patient with CVD, CKD, an patient with psychological disorder etc. ethical clearance was obtained from the ethical committee, Dhaka medical college hospital Dhaka.

Methods:
A hospital based Prospective Randomized double-blind study among adult consented patients aged 18 to 60 years with ASA (American society of Anesthesiologist) class –I and class-ii of both gender. After considering the inclusion and exclusion criteria the patients were randomized to receive Group: A (2mcg/kg oral clonidine) and Group: B (4mcg/kg oral clonidine), one hour (60minutes) before surgery as an oral premedication. All groups were compared for preoperative analgesic, sedation and anxiety level along with changes of heart rate and mean arterial pressure prior to premedication and post operative periods as follows VAS (visual analogue score) pain scores were collected from patients using a standard 10-cm VAS pain ruler VAS measured every hour for first 6 hours, then 4 hours interval for the rest 18 hours and also preoperative VAS taken, pethidine (meperidine) requirement, vital signs (Bp, pulse, mean blood pressure, spo2, respirations), PONV (post operative nausea and vomiting), sedation, recovery score for 1st 6 hours and other side effects (nausea dizziness vomiting) will be measured for up to 24 hours after surgery. Intraoperative analgesic drugs requirement and any postoperative complication were recorded. Detailed demographic data were collected from the informant and recorded in structured case report form. Clinical examination and relevant investigations were done meticulously. All collected questionnaire were checked very carefully to identify the error in the data. The data processing work consist of registration schedules, editing computerization, preparation of dummy table, analyzing and matching of data.

Result:
The present study evaluated the oral premedication with clonidine at 2 microgram per kg or 4 microgram per kg for post operative analgesia and haemodynamic stability of elective upper abdominal surgery patients. A total of 60 patients, 30 in each group, were evaluated. All groups are comparable with respect to the demographic and operational factors. No significant difference were between two groups with respect to age, gender, weight, time between oral premedication to anaesthetic induction, duration of anaesthesia and surgical procedure time.

In our study, we have used oral premedication with clonidine at a dose of 2mcg per kg and 4 mcg per kg and found doses to be effective for post operative analgesic and haemodynamic stability in upper abdominal surgery.
Table 1: Comparison of pre and post operative pain VAS between Group-A (2mcg/kg) and Group B (4mcg/kg).

<table>
<thead>
<tr>
<th>Pain (VAS)(CM)</th>
<th>Group A</th>
<th>Group B</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Just before (preoperative) premedication</td>
<td>0.00</td>
<td>0.00</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>Zero post operative hour</td>
<td>0.00</td>
<td>0.00</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>1st post operative hour</td>
<td>0.3±0.2</td>
<td>0.00</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>2nd post operative hour</td>
<td>0.1±0.5</td>
<td>0.00</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>3rd post operative hour</td>
<td>0.2±0.3</td>
<td>0.00</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>4th post operative hour</td>
<td>0.9±1.6</td>
<td>0.1±0.12</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>5th post operative hour</td>
<td>2.8±2.2</td>
<td>1.7±1.1</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>6th post operative hour</td>
<td>4.3±2.4</td>
<td>1.3±0.8</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>10 th post operative hour</td>
<td>2.3±0.4</td>
<td>1.9±0.6</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>14th post operative hour</td>
<td>1.9±0.7</td>
<td>3.3±1.2</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>18th post operative hour</td>
<td>2.0±0.8</td>
<td>2.3±0.6</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>22nd post operative hour</td>
<td>1.9±1.4</td>
<td>1.3±1.5</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>24th post operative hour</td>
<td>1.6±1.2</td>
<td>1.3±0.2</td>
<td>&gt;0.05</td>
</tr>
</tbody>
</table>

Repeated measure ANOVA statistics was done to analyze the “P” refer to over all statistical difference between two groups S= significant intensity of post operative pain measured on VAS showed Group A expressed highest VAS at 6th post operative hour Group –B showed highest VAS at 14th post operative hour (Table 1). So it is very clear to us that pethedine requirement of Group B (4mcg/kg oral clonidine) is less than Group A (2mcg/kg oral clonidine).

Fig: 0 post operative hours, 1 to 6 is first 6 post operative hours gradually. 7th for 10th post operative hour, 8th for 14th post operative hour, 9th for 18th post operative hour, 10th 22nd post operative hour, 11th for 24th post operative hour

Discussion

This prospective comparative study was carried out with an aim to evaluate the specific and effective analgesic dose of oral clonidine for premedication that plays an important role in post operative analgesia and stabilized the haemodynamic of patient who was operated by upper abdominal incision.

4 mcg per kg body weight of oral clonidine premedication in upper abdominal surgery is more effective than 2mcg per kg body weight, for post operative analgesia and haemodynamic stability, because of its large incision areas need more analgesic. On the other hand laparoscopic cholecystectomy 2mcg per kg is enough for it small incision.

Sung CS et al. 18 found that the post operative analgesic requirement of clonidine premedicant (150 mcg) was less and duration of first dose analgesic was prolonged (4.11 5.65) or 3/5 hours in case of laparoscopic cholecystectomy. In our study group A, which was premedicated by clonidine 2mcg per kg body weight was similar to the above mention study and it was suggested that the improvement of dose in certain level improve the analgesic duration on the post postoperative analgesic requirement period.

Sing S Arora et al. 10 2011 found that the premedicant by oral clonidine 150 mcg was significantly decreased post operative analgesic requirement and improved perioperative haemodynamic stability and a reduction in intraoperative anaesthetic in case of laparoscopic cholecystectomy. They found most of the patient in clonidine group require no meperidine or only one dose during post operative 24 hours period, while more patient in the placebo group required 2 or more dose of meperidine. They suggested that administration of oral clonidine 150mcg as a simple and cost effective form of premedication in patient undergoing laparoscopic surgery. In our study the above mention dose was match with our group A (2mcg/kg). However in case of large upper abdominal surgery dose of clonidine need to improved up to 4 or 5 mcg per kg. Brand JM 19 suggested that the preoperative use of oral clonidine (3.5mcg/kg) followed by intravenous infusion of clonidine post operatively was found improved the haemodynamic profile associated
with anaesthetic discontinuation, thus further proving its anesthetics sparing effect. Mikawa K et al. 1996, 16 commented that oral clonidine premedication reduces post operative pain in children. They had recorded visual analogue score of 29 children to assess the post operative pain after minor surgery. They had given premedication 2mcg/kg one group, 4mcg/kg another group, and an placebo group. They had found VAS for placebo group was (6.6 ±1.7) (mean ± sd, n=11) and VAS for group (2mcg/kg) was (5.7 ±1.5) (mean ± sd, n=10) and group (4 mcg/kg) was (4.4 ±1.6) (mean ± sd n=8). According to their findings it was proved that different dose of clonidine differ in the post operative pain management which is also granted our study that clonidine premedication is dose specific and surgery specific.

Regarding haemodynamic stability at post operative period fist 24 hours most of the patient of group B (premedicated by oral clonidine 4mcg/kg) was stable and in group A (premedicated by oral clonidine 2mcg/kg) also stable except few hours when the analgesic effect of low dose clonidine declined. In the study of Hayash Y et al 1993 they found preoperative oral clonidine (5mcg/kg) decreased the dose of droperidol necessary to maintain the haemodynamic stability at perioperative period in aortic surgery which support our results of group B.

Regarding mean blood pressure changes in post postoperative period there was significant difference found in both groups. Group B (4mcg/kg) was found more effective in stabilizing the blood presser than group A (2mcg/kg). This result is supported by previous studies like kalra NK et al 2011 20 Found that clonidine 1.5 mcg/kg is far better than clonidine 1mcg/kg in suppressing haemodynamic change during pneumoperitonium in lap cholecystectomy. Thalikder HA et al. 2015.8 Also found in lap cholecystectomy, clonidine 150 mcg effectively attenuated the raise of heart rate and mean arterial blood pressure indicating inactivation of catecholamine’s. Borah et al 2017 4 found oral clonidine premedication 1mcg/kg is effective in blunting the haemodynamic response to laryngoscopy in patient who under goes surgery under general anaesthesia , intraoperative and post operative period. Mihossoni et al 2017 6 found in their study clonidine 200mg induced less increase heart rate at 8 hours of operation and MABP peroperatively. Above study closely support our result that 4mcg/kg oral clonidine premedication are more effective to stable the haemodynamic at whole post operative period 24 hours after upper abdominal surgery and low dose 2mg/kg only for operative and early post operative period.

Conclusion
Upper abdominal surgery with large incision area. (like- hepatobiliary surgery, gastrectomy, esophagectomy, hepaticotomy, and whipples operations) causes severe post operative pain, for post operative pain management and to maintain hemodynamic stability, only large dose opioids or opioids with low dose oral clonidine(2mcg/kg) is inadequate. However large opioids doses cases unwanted side effect, respiratory complication, increase hospital stay and morbidity. As a part of multimodal analgesic approach, in upper abdominal surgery, 4mcg/kg oral clonidine premedication is more effective to postoperative pain control and hemodynamic stability than 2mcg/kg oral clonidine premedication.

References:
2. Islam A, Hossain M, Akhtaruzzaman AKM,


