COMPARISON OF EFFICACY OF PLAIN LIGNOCAINE WITH LIGNOCAINE AND CLONIDINE IN INTRAVENOUS REGIONAL ANAESTHESIA FOR UPPER LIMB SURGERY
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ABSTRACT: INTRODUCTION: IVRA is a simple, reliable, and effective technique with rapid onset of action, rapid and prompt recovery after tourniquet release. It provides good analgesia, adequate muscle relaxation, & bloodless operative field. It is widely applicable to patients of different ages and physical status for operations & cost effective. Lignocaine though preferred local anesthetic agent has limitation of short duration of anesthesia & inability to provide postoperative analgesia various additives were added to it. In this study we compared efficacy of clonidine as an adjuvant to lignocaine with plain Lignocaine. MATERIAL & METHOD: Patients undergoing upper arm surgery were included in this study & are divided in two groups. The proximal circulatory isolation of arm was done by placing a pneumatic tourniquet around arm. In group C IVRA was given by 1ug/kg clonidine & 0.5% preservative free lignocaine in a dose of 200mg diluted up to 40 ml & in group L 0.5% preservative free lignocaine 200 mg diluted up to 40 ml. Tourniquet was deflated at least 30 mines after injection of drug. AIMS & OBJECTIVES: To compare onset and quality of sensory analgesia, Onset and quality of motor blockade, onset & severity of tourniquet pain, Complications like hypotension &bradycardia during the procedure, recovery from sensory and motor blockade duration of Postoperative analgesia in group C with group L. CONCLUSION: We observed that using Clonidine in dose of 1 ug/kg as an adjuvant to Lignocaine in IVRA does not have early onset of sensory blockade, increased tourniquet tolerance, delayed tourniquet pain and extended post-operative analgesia .Neither systemic side effects like nausea, bradycardia, hypotension, convulsion nor local complications like hematoma were observed.

INTRODUCTION: Intravenous regional anaesthesia was originally introduced by the German surgeon August K. G. Bier in 1908; thus the name, "Bier block". Dr. Bier described a complete anaesthesia and motor paralysis after intravenous injection of Prilocaine into a previously exsanguinated limb. It is a simple, reliable, and effective technique with rapid onset of action, and prompt recovery after tourniquet release. It provides good analgesia, adequate muscle relaxation, & bloodless operative field, widely applicable to patients of different ages and physical status for operations & cost effective. Poor postoperative analgesia, limited duration of anaesthesia (<90 minutes), the potential for local anesthetic toxicity, nerve damage and compartment syndrome are the disadvantages of intravenous regional anaesthesia.
Various studies are carried out using adjuvant to the local anesthetics solution to improve quality of analgesia in intravenous regional anaesthesia. Adjuvant like Ketamine, Meperidine, Morphine, Sufentanil, Ketorolac, Tramadol, Clonidine, Dexmedetomidine, Magnesium sulphate, Pancuronium, Nitroglycerine has been used. Clonidine is a centrally acting alpha-2 agonist having sedative property. It improves intraoperative stability by reducing catecholamine release and decreases post-operative analgesic requirement. In peripheral nerve blocks, it prolongs the duration of block. Side effects include bradycardia, hypotension, sedation, dry mouth, respiratory depression.

In our study we added 1ug/kg (1ml) Clonidine to preservative free 0.5% Lignocaine 200mg diluted up to 40 ml (Group C) and compared it with 0.5% preservative free lignocaine 200 mg diluted up to 40 ml plus 1ml normal saline (Group L).

AIMS AND OBJECTIVES: This study was conducted in 60 patients undergoing forearm surgeries in orthopedic department. We compared the efficacy of clonidine 1ug/kg added to 0.5 % Lignocaine with 0.5 % Lignocaine for intravenous regional anaesthesia. We compared onset of sensory blockade and motor blockade, intensity of tourniquet pain, intraoperative complications, duration of postoperative analgesia and number of analgesic doses required within first 24hrs in both groups.

MATERIAL & METHOD: After approval of hospital ethical committee, this study was conducted in 60 patients posted for upper limb surgery. Patients were randomized in 2 groups by computer generated randomization with 30 patients in each. Patients with age 15-50 years, ASA grade I and II, weight 50-60kg were included in this study. Patients with sickle cell disease, peripheral vascular disease, hemolytic disease, refusal for regional block, hypersensitivity to Lignocaine or Clonidine, major systemic illness & local infection were excluded from the study. All equipment’s and drugs necessary for resuscitation and general anaesthesia were kept ready. Continuous Electrocardiogram, noninvasive blood pressure, and pulse oximeter were used for monitoring. Pulse rate, mean arterial pressure (MAP) respiratory rate, SpO2 (peripheral O2 saturation) were noted. A tourniquet was placed on the arm to be blocked. We used a “double cuff” tourniquet was used to increase the reliability of the technique and reduce the tourniquet pain.

A 22-gaun intravenous cannula was introduced in the dorsum of the hand to be anesthetized. The arm was elevated for 3-5 minutes to allow passive exsanguination. On the opposite hand a 20 gauge intravenous cannula was put for crystalloid infusion intraoperatively at a rate of 4ml/kg/hr. Patient received intravenous Midazolam 1mg and Fentanyl 100µgm as premedication. Pneumatic double cuff tourniquet was placed around the upper arm, and the proximal circulatory isolation of arm was verified by inspection, absence of radial pulse, loss of pulse oximetry reading in ipsilateral index finger.

In patients from GROUP L 200mg of Lignocaine was diluted with normal saline up to 41 ml & in GROUP C 1ug/kg clenidine and 200mg of Lignocaine was diluted with normal saline up to 40 ml (total volume of 41ml). The solution was injected over 90 seconds in the hand to be operated. Sensory block was assessed by pinprick with 22 G short bevelled needle every 30 seconds. Patient’s response was evaluated in the dermatomal sensory distribution of medial and lateral brachial cutaneous, ulnar (little finger, hypothenar eminence), median (thenar eminence, index finger) and radial (forearm and first web space) nerves.
Sensory block was graded as Gr. 4 - excellent with no pain, Gr. 3 - minor pain with no need of supplemental analgesic, Gr. 2 - moderate pain needed supplemental analgesics (Fentanyl 1µg/kg), Gr. 1 - severe pain (General anaesthesia needed). Motor function was assessed by asking the subject to flex and extend his fingers, wrists and elbow and was graded according to Bromage scale. Time required for motor paralysis was defined as time elapsed from injection of study drug to complete motor block. Motor block was graded as Grade 4 - no movement, Grade 3 - movement only at interphalangeal joint, Grade 2 - movement at interphalangeal and wrist joint, Grade 1 - movement at interphalangeal, wrist and elbow joint.

After sensory and motor blockade, the distal tourniquet was inflated to 100 mmHg above systolic blood pressure, the proximal tourniquet was released, and surgery was started. MAP (mean arterial pressure), HR (heart rate), SpO2 were monitored immediately after distal tourniquet inflation and 5, 10, 15, 20, 30, 40 and 50 min after injection of anaesthetic and after release of tourniquet.

Assessment of tourniquet pain was made by visual analogue scale (0 - no pain and 10 - worst pain imaginable) measured before and after tourniquet application and 5, 10, 15, 20, 30, 40 and 50 min after study drug application. With VAS (visual analogue scale) more than 4, Inj Fentanyl 1µg/kg was given to reduce pain. Changes in heart rate and blood pressure (25% decrease or increase from baseline) were considered significant. Treatment for bradycardia, tachycardia & hypotension and decrease SpO2 if noted was started immediately. The tourniquet was deflated 30 minutes after injection of drug and not kept inflated for more than 90 mints.

At the end of surgery tourniquet was deflated by intermittent cuff deflation re-inflation technique which was repeated for 3 times at 10 seconds interval. Sensory recovery time was noted (time elapsed from tourniquet deflation to recovery of sensation in all dermatomes, determined by pinprick test). Motor block recovery time was noted (time elapsed from tourniquet deflation to movement of fingers, hand and forearm). Postoperative analgesia was assessed every 15 minutes as per VAS (visual analogue scale) in the first hour and later every one hour till score was 4 or more. The first analgesic requirement time was noted (time elapsed from tourniquet release until first patient request for analgesic).

Data was analysed by Student's t tests and Chi square test. The data which included the hemodynamic parameters, SpO2, duration of analgesia were calculated and compared with baseline values within each group using the software ‘GraphpadPrism5’. For comparing quantitative data between the study groups unpaired ‘t’ test was applied. Comparison of non-parametric (qualitative) data between the study groups was done using Chi-square test, Chi-square test for trend and Fisher Exact test depending on types of data. Statistical significance is indicated by conventional symbols: *P<0.05: Statistically significant **P>0.05: Statistically non significant

RESULT: Demographic data for age, sex and weight was comparable in both groups of patients. Onset of sensory blockade was earlier (281± 72.84 seconds) in group C than in group L (382±74.80 seconds). The difference in mean time of onset of sensory block between group L and group C was statistically significant (p<0.001). The difference in mean time of onset of motor block between group L (637.33±115.63 seconds) & group C (579±130.65 seconds) was not significant (p>0.05). Patients in group C had better tolerance of tourniquet than those of group L. No rescue analgesic was required in group C while five patients from group L required Inj. Fentanyl & one patient required supplementation of Inj. Ketamine intraoperatively. Better intraoperative cardiac stability was observed in Group C, as compared to Group L.
Onset of pain after release of tourniquet in group C was significantly prolonged than in Group L. All patients from group L had pain within 2 hours after the release of tourniquet while the patients from group C had pain after 2 hours of release of tourniquet. All patients from group L and only 5 patients in C group required analgesics within first 4 hrs. and patients from group C experienced prolonged analgesia and subjective comfort. Mean time for first analgesic consumption in group L was 99.034±28.11min. and in group C it was 371.66±172.58mints. Better intraoperative cardiac stability was observed in group C as compared to group L.

The mean recovery time of sensory and motor blocks was comparable in both the groups. Patients from group L required earlier postoperative supplementation of analgesics. Out of 30 patients in group C 5(15.66%) patients required analgesics during first 4 hrs, 9(30%) patients required analgesics during first 8 hrs. remaining 13(43.33%) required analgesics after 8 hrs. postoperatively. We did not observe any complications like bradycardia, nausea, vomiting, sedation, convulsions, light headedness, hypotension in either group.

**DISCUSSION:** Adjuvant is added to local anaesthetic solution in intravenous regional anaesthesia to improve quality of block, prolong post-operative analgesia and decreases tourniquet pain. Clonidine, Dexmedetomidine, Fentanyl, Ketorolac, Tramadol, Nitroglycerine, Dexamethasone, Magnesium sulphate, Sufentanil and Pancuronium has been used so far in intravenous regional anaesthesia as an adjuvant. Data from several clinical investigations support the importance of peripheral adrenergic receptors in the maintenance of sympathetically maintained pain. It was supported by the observations like adrenergic blockade with intravenously administered phenolamine, phenoxybenzamine, or prazosin diminishes pain. Intravenous regional anaesthesia with guanethidine depletes peripheral catecholamine and can relieve sympathetically maintained pain, intradermal injection of norepinephrine rekindles sympathetically maintained pain in patients who have previously undergone sympathectomy and topical application of clonidine has been shown to eliminate hyperalgesia only at the site of drug application.

Clonidine is alpha₂-adrenergic agonist that can affect both central and peripheral adrenergic receptors. Clonidine reduces release of norepinephrine from pre-junctional alpha₂-adrenoceptors in the periphery hence reduces sympathetically maintained pain. It was observed that sympathetic neural activity might increase pain associated with skin damage. Clonidine inhibits the release of Norepinephrine from prejunctional α₂-adrenoceptors in the periphery & potentially inhibits neural activity in nociceptive pathways. It acts by selectively blocking conduction of A-δ and C fibres and also causes local vasoconstriction, thereby reducing the vascular uptake of local anaesthetics. Clonidine may produce a peripheral analgesic effect by releasing encephalin-like substances. Perineural administration of the Clonidine is also known to prolong duration of analgesia due to depression of nerve action potentials especially in C fibres. Clonidine has shown greater analgesic potential with minimal adverse effects with a dose of up to 150 µg. Adverse effects of Clonidine are sedation, hypotension, and bradycardia. Clonidine has several clinically desirable effects such as, anxiolysis, anaesthetic sparing and cardiovascular stabilizing property. It reduces intraocular pressure & salivary secretion.

It was observed that maximum hypotensive and sedative effects usually occur with concentrations of clonidine in plasma between 1.5 and 2.0 µg/ml. This might be due to the fact that plasma concentrations of clonidine (0.12 ± 0.05 µg/ml) obtained 30 min after deflation of the tourniquet was significantly lower than those required for a central sympatholytic effect. Higher doses of Clonidine may provide prolonged analgesia at the expense of excessive hypotension and bradycardia.
sedation, bradycardia, and hypotension. This is seen especially when Clonidine is used in a dose > 2 μg/kg. Hence we preferred the dose of Clonidine as 1μg/kg and did not observe any adverse effect.

Gentili M and et al observed that verbal rating scores for tourniquet pain at 15,30, & 45 min for Clonidine group was significantly lower than plain Lignocaine group.³ Our observations match with above findings. Ruben Scott et al compared 1 μg/kg intravenous Clonidine with 1 μg/kg Clonidine added to the intravenous regional anaesthesia solution. They observed that patients in the clonidine group had a significantly longer period of subjective comfort and they didn’t require any analgesics³. We observed decrease in mean blood pressure and heart rate by approximately 15–20%, which was subsided within 2–4 h postoperatively without rescue drug and reduced dose requirements of analgesic drug.

Guanethidine if added to Lignocaine produces prolonged hypotension, apnoea, and angina.⁶ Opiates Fentanyl, Morphine, Sufentanil and Meperidine can be added to Lignocaine but known to produce nausea, vomiting, and dizziness.⁶ To achieve good quality of motor block atracurium or pancuronium was added to Lignocaine but muscle weakness lasts for several hours after use of Atracurium.⁶,⁷

Neil Roy Connellya had assessed the analgesic efficacy of administering Lidocaine and Ketorolac with either a forearm or upper arm tourniquet for outpatient hand surgery.⁸ He found that forearm tourniquet technique results in a longer duration of sensory block and prolonged postoperative analgesia compared with upper arm intravenous regional anaesthesia while using half the doses of both drugs. Ketorolac was known to cause hematomas at the injection site.⁸

Intravenous regional anaesthesia block with bretylium had used but was discontinued because of the high incidence of orthostatic hypotension which prevented the discharge of patients from the outpatient clinic. Bevinagidad Veerappa tried Fentanyl & Tramadol as an adjuvant to Lignocaine for intravenous regional anaesthesia for upper limb surgery& concluded that Tramadol provides significantly longer post-operative analgesia as compared to Fentanyl added to Lignocaine in surgeries of upper limb.⁹

Lignocaine is the most widely used local anaesthetic for intravenous regional anaesthesia due to safe, rapid onset of analgesia, good muscle relaxation, prompt recovery, less cardio toxicity & minimal neurological sequel as compared to other local anaesthetic drugs.⁶,¹⁰ Chlorprocaine produced thrombophlebitis due damage to vascular endothelium leading to pain on administration.⁶ Prilocaine produced methemoglobinemia and its stability cannot be guaranteed if diluted with less incidence of central nervous system toxicity.⁶ Ware used Bupivacaine for intravenous regional anaesthesia IVRA and found that Bupivacaine was associated with several toxic reactions, cardiac arrest in form of ventricular fibrillation.¹¹ Maxmillan et al used Ropivacaine, and found good intraoperative(2hrs) and postoperative analgesia.¹² Articaine produces erythematous rash in the area where drug is injected.⁶

Ketamine was also used as sole agent for intravenous regional anaesthesia. Ketamine was known to produce loss of consciousness, hallucinations and unpleasant psycho mimetic effects.¹³, ⁶, Meperidine was known to develop of dizziness, nausea, pain at the injection site, flushing of face, corneal injection and throbbing headache.¹³,⁶ Hence these drugs are not used routinely for intravenous regional anaesthesia. Magnesium sulphate as an adjuvant produce faster onset of sensory and motor blockade as compare to ketorolac¹⁴. 200µgm Nitroglycerine
added to lignocaine lead to early onset of sensory and motor block with prolonged postoperative analgesia without any adverse effect. 

Success of intravenous regional anaesthesia needs exsanguination of limb at least for 5 min, use of double bladder cuff tourniquet and tourniquet deflation by intermittent cuff deflation re-inflation technique. Maximum allowable tourniquet time for upper limb is 60 to 90 minute. Inflation of tourniquet beyond this period results in complications like tourniquet pain, nerve injury, post-operative oedema of, pressure sores and thrombosis. Use of Lignocaine with Clonidine was useful for early onset of sensory block and prolongation of analgesia.

CONCLUSION: Using Clonidine in dose of 1 µg/kg as an adjuvant to Lignocaine in intravenous regional anaesthesia IVRA has, early onset of sensory blockade, delayed the onset of tourniquet pain, increased tourniquet tolerance, reduced severity of tourniquet pain and extended post-operative analgesia. No effect on onset of motor blockade was observed. No adverse effects like nausea, bradycardia, sedation, hypotension and convulsion or local complications like hematoma were observed. When administered as part of a regional anaesthetic, Clonidine surely prolongs anaesthesia and analgesia in a dose-dependent manner.

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Table 1

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<tr>
<th>sr no.</th>
<th>parameter</th>
<th>group l mean ±sd</th>
<th>group c mean ±sd</th>
<th>p value</th>
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<tr>
<td>1</td>
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<td>35.83±11.92</td>
<td>36.20±14.33</td>
<td>0.9146*</td>
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<td>56.97±5.411</td>
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<td>sex(m:f)</td>
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<td>23:7</td>
<td>0.5675*</td>
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<td>4</td>
<td>onset of sensory block</td>
<td>382±74.80seconds</td>
<td>281±72.84seconds</td>
<td>p&lt;0.001*</td>
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<tr>
<td>5</td>
<td>onset of motor block</td>
<td>637.33±115.63seconds</td>
<td>579±130.65seconds</td>
<td>p&lt;0.05*</td>
</tr>
<tr>
<td>6</td>
<td>duration of surgery</td>
<td>60.36 min</td>
<td>58.01 min</td>
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<tr>
<td>7</td>
<td>tourniquet pain</td>
<td>5 patients needed rescue analgesics</td>
<td>rescue analgesics not needed</td>
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<tr>
<td>8</td>
<td>time of requirement of first analgesic</td>
<td>99.034±28.11 min.</td>
<td>371.66±172.58 min</td>
<td>p&lt;0.001</td>
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<tr>
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<td>intraoperative complications pain</td>
<td>5</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>nausea &amp; vomiting</td>
<td>1</td>
<td>0</td>
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Table II table showing number of patients who required analgesic doses in first 24 hrs

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<tr>
<th>Duration</th>
<th>No of patients required 1st analgesic dose in Group L</th>
<th>No of patients required analgesic dose in Group C</th>
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<tbody>
<tr>
<td>0-4 hrs</td>
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<td>8</td>
</tr>
<tr>
<td>4-8 hrs</td>
<td>-</td>
<td>9</td>
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<tr>
<td>&gt;8 hrs</td>
<td>-</td>
<td>13</td>
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</table>
The difference in mean time of onset of sensory block between group L & group C was found to be significant (p<0.001)

**Graph showing Time of Onset of Sensory Blockade**

- **X-axis**: Time of onset of sensory blockade in seconds.
- **Y-axis**: Number of patients.
GRAPH SHOWING TIME OF ONSET OF MOTOR PARALYSIS

X axis: Time of onset of motor paralysis in Seconds.
Y axis: Number of patients.

The difference in mean time of onset of motor block between group L & group C was found to be non significant (p>0.05)
The tourniquet pain was comparable in both the groups during first 70 min. of surgery. No significant difference found between two groups. (p>0.05)
Graph showing no. of patients who required analgesic tablets in first 24 hrs.

X axis - showing postoperative period in hrs.
Y axis - showing Number of patients, who required post operative analgesic tablets postoperatively.

As from above all patients from group L required analgesia within first 4 hrs as compared to 5 patients in C group. Patients from Clonidine group experienced prolonged analgesia & subjective comfort than Group L.
Above chart shows, better intraoperative stability in respect of heart rate in Group C, as compared to Group L. Heart rate ranged (70-80 beats/min), in group C, while in group L, it was in between (80-90 beats/min).

X-axis shows: Number of patients.
Y-axis shows: Heart rate
GRAPH SHOWING COMPARISON OF MAP BETWEEN TWO GROUPS

PATIENTS FROM GROUP C HAD BETTER INTRAOPERATIVE STABILITY IN RESPECT TO MAP.

X-axis shows: Number of patients.
Y-axis shows: Mean Arterial Pressure (MAP)