Efficacy of Intra-Arterial Lidocaine for Pain Control Resulting from Transarterial Chemoembolization of Hepatocellular Carcinoma

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ABSTRACT

Objective: To assess the efficacy of intra-arterial lidocaine in peri & post-procedural pain control and the dose of narcotic analgesic required in hepatocellular carcinoma patients undergoing transarterial chemoembolization.

Study Design: Comparative prospective study.

Place and Duration of Study: Armed Forces Institute of Radiology and Imaging Rawalpindi, from Jan to Jun 2019.

Methodology: A total of 60 patients included in this study where 42 males and 18 were females, age range 45-85 years who underwent transarterial chemoembolization for hepatocellular carcinoma, were included in the study. patients were equally divided into two groups, group a (30 patients) who underwent transarterial chemoembolization, received 60 mg of intra-arterial lidocaine each and group b (30 patients) who underwent transarterial chemoembolization, intra-arterial lidocaine was substituted with normal saline. Degree of post-procedural pain was assessed using a subjective method (visual analogue scales score) and an objective method (amount of post-procedural analgesics).

Results: Average peri-procedure visual analogue scale score was 5.06 in group A patient versus 7.2 for those in group B patients (p=0.037). Post-procedure visual analogue scale score in the group A was 2.7 ± 0.520 and that for group B was 4.2 ± 0.761 (p=0.025). Mean of total dose of nalbuphine in group A was 4.96 ± 0.764 mg versus 8.3 ± 1.34 mg for patients in group B (p=0.036). Average length of post procedure hospital stay was 0.9 ± 0.203 and 1.41 ± 0.373 days for group A and group B respectively (p=0.002).

Conclusion: Intra-arterial administration of lidocaine before infusing the embolization particles for transarterial chemoembolization in patients with hepatocellular carcinoma is safe and effective in doses as low as 50 mg for reducing peri & post-procedural pain and reducing dosage of narcotic analgesics.

Keywords: Hepatocellular carcinoma (HCC), Intra-arterial, lidocaine, Pain control and pes (post embolization syndrome), Trans arterial chemoembolization (TACE) & visual analogue score (VAS).

INTRODUCTION

Transarterial chemoembolization (TACE) is a well-known technique for the management of unresectable hepatocellular carcinoma. TACE may be used as a neoadjuvant and bridging to resection or orthotopic liver transplantation1. In most institutions TACE is considered as an option when the patient is not a surgical candidate for the treatment of hepatocellular carcinoma (HCC)2-7. It is indicated as a palliative treatment and considered as the first line therapy for intermediate stage hcc according to the recommendation of american association for study of liver diseases based on randomized controlled trials8. TACE is simply administration of cytotoxic drugs with or without lipiodol, by means of a catheter positioned in the tumor supplying hepatic artery followed by the administration of embolizing agents such as spherical gelatin or polyvinyl alcohol particles9. In most patients TACE commonly causes procedure-related abdominal pain either during or after the procedure, sometimes even in patients who did not experience pain during the procedure. The pain is severe enough to necessitate narcotic analgesia. Post embolization syndrome (PES) is a common complication after embolic procedures and it is a frequent cause of extended inpatient hospital admissions. PES is a self-limited constellation of symptoms consisting of fever, unremitting nausea, general malaise, loss of appetite, and variable abdominal pain following the procedure. Although a definite cause is unknown, this syndrome is thought to be a result of therapeutic cytotoxicity, tumor ischemia and resulting intrahepatic and extrahepatic inflammation10.

METHODOLOGY

This comparative prospective study was conducted at armed forces institute of radiology and imaging (AFIRI) Rawalpindi. It was approved by institutional ethics committee-IERB approval certificate number is 00012, prior to the procedure informed consent was obtained from all of the patients included in study.

The sample size calculated was 60 patients with hepatocellular carcinoma using the who sample size
Intra-Arterial Lidocaine

calculator, keeping the confidence level 1.96 (95%), margin of error 0.49, baseline level of indicators 0.5, design effect 1.5, expected response rate 0.8 and number of age/sex estimates. Simple random sampling technique was used in this study.

Inclusion criteria for TACE were corrected coagulopathy, single or multifocal HCC, tumor volume <50% of liver, patients with child A or B, no main portal vein tumor thrombus and no extrahepatic disease.

Inclusion criteria for administration of lidocaine were same as that of TACE.

Exclusion criteria of TACE were uncorrected coagulopathy, infiltrative type of HCC, tumor volume >50% of the liver, patients with child-C according to child pugh classification, extra hepatic metastasis of HCC that was confirmed by CT chest, abdomen and pelvis and thrombosis of main portal vein.

Exclusion criteria for administration of intra-arteria lidocaine were patients on pre-existing regular analgesics, patients who required conscious sedation and any contra-indication to lidocaine like heart block.

All patients were subjected to standard evaluation for TACE in HCC. The diagnosis was made either by typical imaging criteria of hcc with triphasic dynamic contrast studies by CT (fig-1) or MRI in addition to serum level of alfa fetoprotein or in equivocal cases by of histopathology. Pre-procedural standard investigations for TACE to assess hepatic and renal function and coagulation status were done.

Continuous monitoring of blood pressure, pulse and ECG was done during the entire procedure. TACE was performed after visceral angiography to evaluate arterial supply of the HCC and evaluate patency of portal vein. HCC arterial supply was accessed by selective catheterization using standard 5 fr catheter or 2.8 fr coaxial technique using progreat microcatheter (terumo). once the catheter was in suitable position, a slow injection of doxorubicin 50mg mixed with 10ml of lipiodol (ultra-fluids guerbet france) thus forming total volume of chemotherapeutic emulsion was about 15 ml was administered (fig-2). In some cases, deb-TACE was performed in which 50 mg doxorubicin was mixed with 25 mg hepsphere microsphere 30-60 microns

Figure-1: a) Contrast enhanced triphasic CT liver showed segment VIII lesion showing arterial phase enhancement, b) and washout on portal venous phase: Consistan with hepatocellular carcinoma.

Figure-2: a) Super selective angiogram (with microcatheter) of segment VIII HCC supplying branch of right hepatic artery showed tumor blush, b) post conventional TACE super selective angiogram of segment VIII HCC supplying branch of right hepatic artery with microcathetr showed satisfactory flow stasis in tumor feeding artery and good lipoidal retention in tumor bed.
(merit medical) mixed with 10ml of non-ionic contrast-ultravist (bayer) againmaking the total volume of 15ml. in a few cases, only embolization was done with pva particles (45-150 and 255-350 microns) merit medical) mixed with 10ml of non-ionic contrast-ultravist (bayer) again making the total volume of 15ml for embolization only. The chemotherapeutic emulsion was infused under fluoro guidance. Patients were randomly divided into two groups, 3ml of lidocaine 2% (60 mg) was infused in group a just before the infusion of chemotherapeutic emulsion or pva particles and normal saline was infused in (group B) patients who did not receive intra-arterial lidocaine. In both groups, the procedure was concluded by infusing aliquots of polyvinyl alcohol particle (PVA) size of 45-150 micron till satisfactory flow stasis in tumor supplying hepatic artery.

Good hydration was assured for patients before and after procedure by iv normal saline infusion till the ability to drink.

Pain score was recorded using visual analog scale (vas) considering 1 as minimal discomfort and 10 as the most severe pain (fig-3). Pain scores were recorded 4 times on the procedure day and then two times per day. On the procedure, first recording was at the time of infusion of chemotherapy and embolizing material, second was immediately after the procedure, third reading was after 2 hours and last reading was in the evening about 6 hours after the procedure. Similar scores were taken twice a day for next two days, at 6am and 6pm. Pain scores of ≥5 was considered as significant pain that required analgesia. The scores were entered in the prescribed proforma. Wherever necessary, post procedure analgesia was provided by intravenous nalbuphine and both the mean dose and total dose of the drug were recorded. Hospital stay was also recorded for each patient in both groups. comparison of mean of the vas scores, requirement of nalbuphine and hospital stay between the two groups was done. Other symptomatic medication like ondansetron hydrochloride (onset) 4 mg slowly intravenous infusion for nausia was given. Statistical analyses were performed with the independent sample t-test using SPSS. A p-value <0.05 was considered significant.

RESULTS

Sixty transarterial chemoembolization (TACE) procedures were carried out for 60 consecutive patients where 42 (70%) patients were male and 18 (30%) were female. Age of the patients ranged from 45 years to 85 years, mean age was 60.4 ± 7.63 years. Lidocaine group consisted of 30 TACE procedures and every one of these patients received 60 mg intra-arterial lidocaine during chemoembolization. Placebo group consists of 30 TACE procedures in 30 patients in whom intra-arterial lidocaine was substituted with normal saline solution. The patient demographic criteria, child pugh score, tumor size and doses of chemotherapeutic emulsion and amount of pva particles used were comparable without statistically significant difference. TACE technique was selective & super selective or segmental and lobar for both lidocaine and placebo groups. The majority of patients had right lobe lesions. there was no recorded inadvertent embolization of the gall bladder supplying artery. There were no vascular complications during TACE procedure such as dissection or spasm of hepatic artery. The infused dose of chemotherapeutic emulsion till tumor bed saturation ranged from 0.5-1.0 of the prepared chemotherapeutic emulsion for the lidocaine group and from 0.5-0.9 of the emulsion in the placebo group, the dose difference was not statistically significant and so did the volume of the emobilization particles (PVA). There were no recorded changes in blood pressure or arrhythmias in patients who received lidocaine. Moderate inter procedure pain was noticed in the entire lidocaine group (n=30) with vas (visual analog scale) score of 4-6 compared to 5 (16.6%) in the placebo group. Severe pain was noticed in 25 (83.3%) of placebo group. The average periprocedural pain score was less in the lidocaine group than in the placebo group. The average vas was 5.06 versus 7.2 for lidocaine and placebo groups respectively (p=0.037) (table). Post procedure pain was significantly lower in lidocaine group versus placebo group. vas score for pain in the lidocaine group was 2.7 ± 0.520 and that for the placebo group was 4.2 ± 0.761, the difference is significant (p=0.025). Vas score for pain on second day in the lidocaine group was 1.73 ± 0.583 and that for the placebo group was 3.10 ± 0.844.
the difference is significant \((p=0.018)\). The frequency of analgesic demands was higher in patients who did not receive lidocaine and total dose of nalbuphine in the lidocaine group was 4.96 ± 0.764 mg versus 8.3 ± 1.34 mg for patients in the placebo group \((p=0.036)\) (table).

**Table: Difference between lidocaine and placebo group as regards to pain score, nalbuphine requirement and length of hospital stay.**

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Nausea was encountered in 27 (90%) of patients belonging to the lidocaine group and it was 100% in those belonging to the placebo group. Incidence of post procedure fever was not statistically different in lidocaine and placebo groups, 4mg of ondansetron hydrochloride (onset), was slowly given as intravenous infusion to patients in 27 (90%) of TACE procedures in the lidocaine group and was given to the entire placebo group patients. The average length of post procedure hospital stay was 0.9 ± 0.203 day and 1.41 ± 0.373 days for lidocaine and placebo groups respectively. that difference was statistically significant \((p=0.002)\).

**DISCUSSION**

Post embolization syndrome (PES) is experienced after TACE procedures in 80-90% of patients. It has widely variable manifestations but often includes pain, fever, nausea and vomiting. PES can last from a few hours to a few days 1. Although it is self-limiting condition, it is a major complication of hepatic TACE causing longer hospital stay9. As the large doses of intravenous narcotic analgesics needed to control the pain leads to altered mental status and respiratory depression; therefore intensive monitoring is required. Narcotic analgesics also potentiate severe post embolization nausea and vomiting11. The exact explanation of pain component of pes in TACE is not known but many hypothesis are postulated such as ischemia and transient swelling of liver parenchyma that stretches the liver capsule or accidental embolization of the arterial supply of gall bladder. Severe pain during the procedure can be explained by irritant effect of the chemotherapeutic emulsion on the hepatic artery branches11,12. Direct irritation of the arterial wall by the chemotherapeutic emulsion is one of the theories of pain component of pes. Daniel et al found that post TACE pain was lower after the first session and this could be explained by lower dose of chemotherapy in the successive TACE sessions than that of the first TACE session13. In super selective TACE, the arterial system which is in contact with irritant chemotherapeutic emulsion is at minimum & the dose is lower than lobar TACE technique and carries less risk for non-target embolization of the gall bladder. Inadvertently emobilized gall bladder artery was considered as one of the theories of pain component of pes. Contrary to daniel et al, patel et al found that repetition of TACE is not a predictor of pain component of pes as their hypothesis was that: ischemic pain was the main mechanism and vascular irritation by chemotherapeutic emulsion was not the major cause of pain14. In a study by coldwell et al excellent analgesia during hepatic TACE was achieved with a celiac plexus block. However, this method seems to be risky and time-consuming15. Intra-arterial lidocaine administration during TACE has been known not only for reduction of severity of the pain that is associated with TACE but facilitates faster recovery as well11,12. Lidocaine has been shown to help control the painful response to the injection of iodinated contrast material in peripheral arteries16,17. Molgaard et al11 studied the use of intra-arterial lidocaine in hepatic arterial branches prior to and during TACE. This resulted in a significant decrease in the amount of morphine (narcotic analogue) required during the procedure as well as the need for subsequent postprocedure morphine drip. It remains clear that the sequelae of pain during and after TACE such as shallow respirations and paralytic ileus can complicate patient management. Therefore intra-arterial lidocaine administration is recommended because it is much easier and less time-consuming method than celiac plexus block. The mechanism of the analgesic effect of intra-arterial lidocaine in hepatic TACE is unclear. Hartnell et al12 suggested that lidocaine has a direct local effect after diffusion into the arterial wall & liver parenchyma and this effect will be prolonged in tumors where blood flow is occluded preventing washout of the agent. Lee et al found that transcatheter administration of lidocaine immediately before infusion of chemotherapy had significantly better effect on pain control than after chemotherapy emulsion13. Lee et al found that patients who had received lidocaine by intra-arterial route during TACE procedure needed smaller doses of narcotic

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analgesics than those who had not received lidocaine\textsuperscript{18}. Different protocols for intra-arterial lidocaine administration were applied. Sharma \textit{et al}\textsuperscript{19} used lidocaine intermittently during the TACE procedure. Hartnell \textit{et al}\textsuperscript{12} injected lidocaine at varying intervals before and during TACE up to 4 times. The dose of lidocaine used in their study (maximum 105mg injected over 10-20 min) was safe and effective. However Lee \textit{et al}\textsuperscript{18} concluded that intra-arterial administration of lidocaine just before TACE was much useful than after TACE as regards to pain control and post procedure requirement for narcotic.

In our study we used lidocaine as bolus rather than infusion; we found that 60 mg of lidocaine was effective to alleviate pain during the procedure and reduced pain score and analgesic dose after the procedure. The post procedure pain control was significant in terms of low pain scores and smaller doses of analgesic requirement.

Superselective embolization is better than lobar TACE as regards to control of non-target embolization especially inadvertent embolization of the gall bladder which in some theory is the main cause of pain component of PES\textsuperscript{13}. Super selective TACE reduces the number and length of arteries exposed to irritant chemotherapeutic emulsion which is one of the theories of pain component of PES. Super selective TACE is associated with reduction in the chemotherapeutic dose infused which is another advantage of superselective over lobar TACE technique in addition to much better tumor necrosis \textsuperscript{20}. Lidocaine is metabolized by the liver and its half-life is about 2 and 2.5 hours with normal liver functions. Lidocaine metabolites and unchanged drug are excreted by the kidneys. Because of the rapid rate at which lidocaine is metabolized, any condition that affects liver function may alter lidocaine kinetics. The half-life may be prolonged two fold or more in patients with liver dysfunction such as cirrhotic patient with HCC. Entrapment of lidocaine in the vascular bed of liver tumor that is partially saturated by chemotherapeutic emulsion and infused embolization particle may prolong the duration of action of lidocaine but this cannot explain the extended duration of action for following few days\textsuperscript{21}. Lidocaine has a potential anti-inflammatory effect\textsuperscript{22} however; there is still a lack of well-designed studies to support this hypothesis. Interestingly, kogut \textit{et al}\textsuperscript{23} found that prophylactic intra-arterial administration of steroids in TACE procedures did not affect analgesic agent use and had a minor effect on antiemetic requirements.

In our study, although pre-procedure intra-arterial lidocaine administration improved pain component of pes, the length of hospital stay was slightly different in patients who received lidocaine versus patients who received placebo. Our results agreed with those of other authors\textsuperscript{24}. The limitation of our study was that the number of patients and procedures were slightly small.

CONCLUSION

Intra-arterial administration of lidocaine just before infusion of embolization particles in doses as low as 60 mg is sufficient for pain control during TACE procedure and helps in pain control after the procedure. Furthermore in order to reduce the incidence of post-procedural pain and dose of post-procedural analgesics, we recommend routine pre-TACE administration of lidocaine because post-procedural pain might develop even in patients who did not feel any pain during the TACE.

CONFLICT OF INTEREST

This study has no conflict of interest to be declared by any author.

REFERENCES