EVALUATION OF CHONDROPROTECTIVE EFFECTS OF INTRA ARTICULAR PIROXICAM IN POST-TRAUMATIC OSTEARTHRITIS MODEL OF RAT

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ABSTRACT

Objective: To evaluate the chondroprotective effect of piroxicam in post-traumatic osteoarthritis model of rat.
Study Design: Laboratory based experimental study.
Place and Duration of Study: Pharmacology department, Army Medical College, Rawalpindi, from Apr to Jun 2019.
Methodology: Project included sixteen rats of Sprague Dawley breed. Osteoarthritis was induced in anesthetized rats by surgical removal of medial meniscus and anterior cruciate ligament resection. After that rats were randomly allocated in two groups with eight rats in each group. Rats of group I were positive control that received 0.2 ml saline intra articularly once weekly for four weeks. Meanwhile rats of group II (treatment group) received 50 µl piroxicam intra articularly once weekly for four weeks. One week after the drug intervention, radiograph of the right knee joint of all rats were taken. Animals were then sacrificed with inhaled chloroform and part of proximal tibia was obtained for histopathological analysis.
Results: Comparison of radiographs of both groups depicted a significant p-value of <0.01. Meanwhile mean histopathological score of control group and treatment group were 11.50 ± 1.195 and 6.50 ± 1.195 respectively with a p-value of <0.01.
Conclusion: Intra articular administration of piroxicam in post-traumatic Osteoarthritis model of rats resulted in improvement in radiographic grades and histopathology scores.

Keywords: Chondroprotective effects, Osteoarthritis, Piroxicam.

INTRODUCTION

Osteoarthritis (OA) was one of the conspicuous causes of mobility-related disability. Age related degenerative changes are the foremost cause of development of OA in older people. Meanwhile trauma to affected joint is the leading cause of development of OA in people younger than 45 years of age1. Young person who sustains a joint injury are recognized to be at significantly increased risk of developing OA as compared to non-injured one. OA that progresses after joint injury is termed as post-traumatic OA (PTOA). Individuals with PTOA approximately accounts 12% of all cases symptomatic OA2. Risk of PTOA is more common in younger healthier and more active people as compared to idiopathic OA which is common in old age. That’s why patient population suffering from PTOA requires medical and surgical intervention at a much earlier age in life. Knee joint are the the most vulnerable joint for developing PTOA3. Any injury severe enough to cause anterior cruciate ligament (ACL) tear or meniscus destabilization frequently leads to PTOA in knee joint4. PTOA affects whole joint architecture including bone, ligaments, bursas, synovium and periarticular muscles. Non pharmacological measures that use to manage PTOA is awareness and education of patient about disease pathology and to persuade him to avoid triggering factors of disease progression5. Meanwhile drug groups are used to lessen the severity of symptoms and to delay the progression of disease. Nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids and viscosupplement substances are widely used drug groups in the treatment of PTOA6. Piroxicam is one of the investigational NSAID that is commonly prescribed to patients of PTOA. It not only relieves pain but also subsides inflammatory component of PTOA. Long duration of action is the advantage of Piroxicam to many other NSAIDs. Piroxicam is a non selective reversible inhibitor of COX enzyme. It competes with arachidonic acid to bind with COX enzyme. Inhibition of COX enzyme leads to impaired generation of prostaglandins (PGs), prostacyclins and thromboxanes. COX I enzyme is present mainly in the stomach while Prime location of COX II is the site of inflammation and tissue injury. Inhibition of COX I causes disruption of physiological process of gastric protection that leads to GI symptoms ranging from mild dyspepsia to erosion and rarely severe bleeding. Similarly inhibition of COX II leads to decrease concentration of PGE1 and PGF2α in inflamed tissues that not only relieves pain but also lessens inflammation.
Chondroprotective Effects of Intra Articular Piroxicam

To avoid gastrointestinal symptoms of oral therapy as well as to obtain targeted effects of the drug, Intra articular (IA) route of drug administration is practiced. IA administration of therapeutic agents has been clinically explored to selectively deliver active compounds to their site of action in the treatment of osteoarthritis, rheumatoid arthritis, and joint pain. Direct IA delivery of active compounds to affected tissues offers the chance to boost therapeutic outcomes with lower dose, while reducing systemic exposure and undesirable adverse effects. Beside COX inhibition, Piroxicam reduces the vasodilation property of vessels in response to bradykinin and histamine. Thus it also reverses the vasodilation of inflammatory tissues. It also lessens lymphokines production from T lymphocytes. Recent studies proposed that Piroxicam is also involved in inhibition of central component of pain mediation. It blocks the B2 receptor mediated pathway in the sensory neurons and thus inhibits bradykinin induced nociception7,8.

Rationale of this animal study was as Piroxicam is an investigational drug in the management of PTOA, we planned this project to detect chondroprotective effects of IA piroxicam in a rat model of PTOA.

METHODOLOGY

It was a laboratory based experimental study that was carried out in department of Pharmacology and Therapeutics, Army Medical College (AMC), Rawalpindi in collaboration with National Institute of Health (NIH), Islamabad. This study was ethically approved from ethics review committee of “Centre for Research in Experimental and Applied Medicine (CREAM)”, AMC. Intervention protocol of this study was about two months from April 2019 to June 2019. Rats were kept and nurtured in animal house of NIH during the complete study period. Preliminary sixteen (16) adult male or non-pregnant female rats of Sprague Dawley breed, of about 02 months and weighing about 400 gram were selected through non probability consecutive sampling. Conventional cages with wood chip bedding were used and animals were identified by a cage label. Environment of rats was maintained with temperature ranged 25 ± 5°C and 12 hours day night cycle. Free excess to clean drinking water and standard rodent diet adlibitum were provided during the whole study period9. They were allocated randomly in two (02) groups with eight (08) rats in each group. Surgical procedure was performed to induce changes similar to PTOA in right knee joint of all rats. They were anesthetized with intraperitoneal injection of 5% xylazine and 1% ketamine10. Skin of the joint was shaved with sterile razor and disinfected with pyodine followed by a para patellar incision on medial side of the joint. Skin and fascias were retracted for the complete exposure of joint. After identification anterior cruciate ligament was removed and medial meniscus was transected. Skin was closed with surgical stapler thereafter. Later on Animals were permitted to move freely in the cage for two weeks11. Then intra articular drugs were administered in the right knee joint of the rats. Rats of control group and treatment groups were injected with 0.2 ml of Normal saline and 70 µl (1.4 mg/ml) piroxicam once weekly for 04 weeks12. We delayed for one week more and then radiographs of corresponding joint of anesthetized (with 5% xylazine and 1% ketamine) rats were taken and graded according to Kellegren and Lawrence grading system with the help of radiologist13. After that animals were sacrificed with inhaled chloroform in a desiccator and sample of proximal tibia was obtained by using 5.5°angled bone cutter14. After histological slide preparation, these slides were scored using Modified Mankin slide preparation15. Obtained data was statistically analyzed using SPSS version 25. Radiographic grades were qualitative parameter and it was compared with Fischer Exact test while histopathological score was quantitative parameters and compared through student t-test. The differences between two observations were considered statistically significant if the p-value was equal or less than 0.05 (p≤0.05).

RESULTS

The rats remained alive healthy and active throughout the study period. One week after the last IA injection Radiographs of right joint of anesthetized rats were taken and graded with the help of radiologist. Radiographs of 02 rats of control group had features of large osteophytes, marked joint space narrowing (JSN) and severe sclerosis and their grade was 04. Meanwhile 04 rats of this group graded with grade 03 because multiple osteophytes, definite JSN and visible sclerosis were features of their radiographs. Two rats

| Groups                          | Grades   | p-value |
|---------------------------------|----------|---------|
|                                 | Grade 0  | Grade 1 | Grade 2 | Grade 3 | Grade 4 | <0.01 |
| Group I (control group)         | -        | -       | 2 (25%) | 4 (50%) | 2 (25%) |
| Group II (piroxicam group)      | -        | 4 (50%) |         |         |         |       |

Table-I: Kellegren and Lawrence grading of rat model of Post traumatic osteoarthritis.
of control group depicted definite osteophytes with possible JSN and grade of their radiograph was 2. Fig-1 is an X-ray of a rat of control group with grade 03. Meanwhile radiograph of half rats of piroxicam group showed characteristics of doubtful JSN and possible osteophyte lipping and their grade was 01 and radiographs of other half of rats of piroxicam group exhibited features of definite osteophytes and possible JSN and their grade was 02. Figure-2 was a photomicrograph of rat of piroxicam group with minimal changes of OA and its grade was 01. Table-I is a cross tabulation of radiographic grades of both group. When these radiographs were compared with fischer exact test, we found significant p-value of <0.01 that explained that intra articular piroxicam exhibited chondroprotective effects in PTOA model of rat at radiological level.

Histopathological analysis of all slides of both group was done under X100 and X400 lens. These histopathological changes were scored according to Modified Mankin scoring system.

Score of slides of control group was in range of 10-13. Six out of eight slides showed marked while rest of two showed mild irregularity in perichondrium. One slide had marked, six had moderate and one had mild fibrosis of perichondrium. Six slides had moderate to mark while two slides had mild to moderate irregularity of organization. All slides showed moderate to marked increase in cellularity of chondrocytes. Five slides showed moderately increased while three slides showed mild increase in chondrocyte clusters. Five slides showed 10-20% while three slides showed 20-80% necrosis of chondrocytes. Fibrinoid degeneration was feature of all the slides of disease control group. Figure-3 is photomicrograph of slide of rat of this group with 13 score. Mean score of group I was 11.50 ± 1.195.

Slides of group II (piroxicam group) scored around 5-8. Mild irregularity of perichondrium was the feature of all slides. One had none, six had mild, while one had moderate fibrosis of perichondrium. Seven had mild to moderate while one had moderate to marked irregularity in organization. One slide showed no, four slides showed mild and three slides showed moderate to marked increase in cellularity of chondrocytes. Seven slides showed mild while one slide showed moderate increase in chondrocyte cluster. 10-20% necrosis of chondrocyte was the feature of all slides. None of the slides exhibited fibrinoid degeneration. Figure-4 is a photomicrograph of slide of rat of piroxicam group with 07 score. It showed features of mild perichondrium fibrosis, moderate irregularity in organization, hypercellularity and 10-20% of chondrocytes necrosis. Mean score of this group was 6.50 ± 1.195. When the histological score of both groups were compared with student t-test, we found a significant p-value of <0.01.

In short we observed a visible decrease of Kellegren and Lawrence radiographic grades and Mean Modified Mankin score of piroxicam group as compared to control group. Inter group comparison of radio-
graphic grades and histopathological score exhibited the p-value <0.01 both times that confirmed the chondroprotective efficacy of piroxicam in rat model of PTOA.

**DISCUSSION**

Nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids and viscosupplement substances are group of drugs that are frequently used in the management of post-traumatic osteoarthritis (PTOA)\(^1\). Piroxicam is an oxicam derivative NSAID that is frequently prescribed in pain conditions including PTOA all over the world. This animal study was considered to find out the chondroprotective effects of intra articular (IA) piroxicam in PTOA rat model. Human admissible dosage of piroxicam that was proficiently effective in rat model was selected by substantial search and literature review. After intervention protocol, when radiographic grades and modified mankin score were compared, differences between rats of piroxicam group and disease control group were statistically significant that confirmed the chondroprotective effects of piroxicam. A similar results found in studies of Park and his colleagues. It has proven in their study that there was a statistically significant difference \((p<0.05)\) of joint swelling and PGE2 level in IA piroxicam treated rats as compared to IA saline treated rat models of OA\(^1\). Likewise research work of Aziza and Hana revealed that IM therapy of piroxicam reduces joint edema and arthritic index statistically significantly \((p<0.05)\) in Freud Adjuvant induced arthritis models of rat\(^1\). Their findings also strengthen our results regarding chondroprotective efficacy of piroxicam. Meanwhile Research work of ljaz ul Haq and his colleagues also proved chondroprotective efficacy of piroxicam \((p<0.05)\) in rabbit models of OA that also reinforce our results\(^1\).

Tenoxicam, an oxicam drug just like piroxicam, used intra articularly in the treatment of surgically induced PTOA rat model. Histological analysis depicted a significant p-value of 0.028 when compared Tenoxicam treated rats with non treated rats. Thus it was claimed that Tenoxicam has chondroprotective effects also signifying our target drug piroxicam might have same chondroprotective effects against OA\(^2\). Likewise meloxicam, another oxicam derivative sister drug of piroxicam, is used in chemically induced rat model of OA. It significantly decreased \((p=0.02)\) heat stimulation of affected joint as well as TNF-α level as compared to control group that also strengthened our results\(^2\).

This study also has some curbs. Though IA route of piroxicam administration leads to less systemic exposure, we did not study the adverse effects. Future studies should be explored that also observe the systemic adverse effects. Similarly a study alike to this contemporary study should be accompanied with prolonged duration between surgical trauma and drug administration. Delay after trauma of up to four to eight weeks may lead to more severe form of disease\(^2\).  

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**Disclosure**

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**CONCLUSION**

IA administration of 70 µl (2mg/ml) piroxicam once weekly for 04 weeks lead to reduced radiographic grades and mean Modified Mankin score of histopathology as compared to rats treated with saline only. Thus we concluded that piroxicam exhibits chondroprotective efficacy in PTOA models of rats.

**CONFLICT OF INTEREST**

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

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