INVESTIGATION OF ANTI-PYRETIC ACTIVITY OF CINNAMON OIL IN WISTAR RAT

Gaurava Srivastav1, Dakshina Gupta2, Anubhav Dubey3

1Research Scholar, Department of Pharmacology Advance Institute of Biotech and Paramedical sciences Kanpur (U.P.) – India.
2Assistant Professor, Department of Pharmacology, Advance Institute of Biotech and Paramedical Sciences Kanpur (U.P.) – India.
3Assistant Professor, Department of Pharmacology, Maharana Pratap College of Pharmacy Kanpur (U.P.) – India.

ABSTRACT

The current exploration work was to explore the antipyretic movement of cinnamon oil (25 Milligram/ kilogram or 50 Milligram/ kilogram p.o). The antipyretic movement was assessed against Brewer's yeast and lipopolysaccharides instigated pyrexia in rodents. In case of lipopolysaccharides instigated pyrexia (50 mg/kg, p.o) is more effective comparison to 25 Milligram/ kilogram. But in case of Brewer's yeast induced pyrexia both low and high dose is more effective of cinnamon oil (25 mg/kg or 50 Milligram/ kilogram p.o) suppress anal temperature significantly 96.42 ± 0.009 and 95.40 ± 0.000 respectively in Brewer’s yeast instigated pyrexia. The exploration result shows that cinnamon oil (25 Milligram/ kilogram or 50 Milligram/ kilogram, p.o) is having a huge enemy of antipyretic potential.

Keywords: Lipopolysaccharides, Brewer's yeast, Cinnamon oil, Pyrexia.

I. INTRODUCTION

Plants play a special position on the globe because they are the source of inspiration for human lifestyles. They are a very well food makers in the world. Plants provide 90 percent of human calorific ingestion and eighty percent of protein intake all at once. Herbs have been utilized as a reliable origin of medicament since the dawn of humanity. India has a vast variety of medicinal herbs, and the people there prefer to use traditional methods of treatment. As a result, it's critical that this magnificent herbal asset be enhanced and employed in accordance with the advancement of development and work wants[1].

Body temperature among 37.22°C and 40.57°C onward is referred to as pyrexia at the same time as upward thrust of frame temperature above 41.66°C is hyperpyrexia. Body temperature rises because of derangement of warmth regulating mechanism. Toxins (pyrogens) act on WBC and bring endogenous pyrogen. This acts at once on anterior hypothalamus and the frame temperature is elevated. Fever takes place because of any of the reasons which include infections (e.g. pneumonia, typhoid fever etc), harm to fearful centres, dehydration, tissue destruction, management of a few tablets etc[2]. Puerperal warmth, Puerperal fever is described because the existence of a heat in a mom extra than and identical to 38°C within side the first fourteen days after giving birth. There are many reasons of any such warmth, however within side the day’s previous to antibiotics, it turned into a signal which turned into very lots dreaded because it had a totally negative prognosis. These days, with set off reputation and remedy of the underlying cause, the final results is substantially better.
The everlasting tree of tropical medicine, cinnamon (Cinnamomum zeylanicum or Cinnamon cassia), is a member of the Lauraceae family. Cinnamon is amongst the most important spices used by people around the world on a daily basis. Cinnamon consists primarily of essential oils and derivatives such as cinna-maldehyde, cinnamic acid, and cinnamate. Cinnamon, in addition to being an antioxidant, anti-inflammatory, antidiabetic, antibacterial, anticancer, lipid-lowering, and cardiovascular-disease-lowering chemical, has also been linked to the prevention of neurological disorders such as Parkinson’s and Alzheimer’s. Cinnamon is employed in the scent and essence sectors and for its fragrance, that may be incorporated into a variety of foods, perfumes, and pharmaceutical items. The most important components in cinnamon are cinna-maldehyde and trans-cinnamaldehyde (Cin), which are naturally present in the essential oil and contribute to the fragrance and a variety of organic properties.

According to a study on Cinnamomum osmophloeum (C. osmophloeum), the essential oil form cinnamon leaves has an inordinate number of Cinnamon. As a result, C. osmophloeum is also utilised as a spice substitute for C. cassia. (E)-cinnamaldehyde, amongst the most essential aspects in crucial oil produced from C. zeylanicum, exhibits an antityrosinase effect, with cinnamaldehyde being the key component responsible for this effect. Procyanidins and catechins are found in cinnamon bark.
II. MATERIALS AND METHODS

Experimental Rodents

Wistar albino rodents of either sex weighing between 150-200g were used for this study. They were procured in the AIBPS animal house Kanpur recognized by the Institutional Animal Ethics Committee (IAEC). Polypropylene limits were used to house (3 for each pen) theanimal at a temperature of 28 ±50C and 12 h light /dull cycle. Hindustan Lever chow pellets were used to feed the animal and water not basic. The animals were kept fasting medium –term going before the examination and this study was approved by IAEC for animal studies (1122/PO/Re/S/2007/CPCSEA) include all framework used in the research.

Chemicals and drugs

Cinnamon oil was gotten from Micro Labs Limited and Paracetamol Injection (Neon Laboratories Limited Andheri East Mumbai Batch No.SM 91145); Yeast concentrates powder (Molychem India Limited Mumbai Maharashtra Batch No.MCR-18566) and Lipopolysaccharide solution sigma Aldrich, Indomethacin Batch No.JIC-013 was gotten from Jagsonal Pharmaceutical Ltd.

III. STUDY DESIGN

1 Brewer’s Yeast Instigated Pyrexia in Wistar Rat

Principle

The subcutaneous implantation of Brewer's yeast blend is known to move fever in Wistar rodents. A decline in temperature can be ended by the association of blends in with antipyretic activity.

Technique

1. 20-Wistar rodents (150-200 gm) of either sex were arbitrarily partitioned into four gatherings.
2. Control Group’s – One percent Tween 80 percent in normal saline solution (o.p.)
3. Standard Group’s – Paracetamol 100 mg/kg
4. Test Group’s – I Cinnamon Oil 25 mg/kg orally
5. Test Group’s – II Cinnamon Oil 50 mg/kg orally

15 percent W/v suspension’s of Brewer's liquefaction in refined water at a portion of 10 milliliter/kilogram body weight was infused in the back below the nape of Wistar rodents. Before experimentation, the rectal degree of heat of all animal groups was recorded utilizing a clinical digital thermometer’s. After 18hr Brewer's yeast injection increase the rectal degree of heat was noted and only creatures showing an influence in the degree of heat of at least 0.60° C or (1° F) were chosen for the study. Control group creatures have given 1% Tween 80% in typical saline arrangement orally and Standard gathering creatures have given Paracetamol drug 100 Milligram/ kilogram, Test bunch I creatures given Cinnamon Oil 25 mg/kg orally and Test gathering – II given Cinnamon oil 50 Milligram/ kilogram orally. After the treatment, the level of warmth of the considerable number of rodents in each gathering was noted at 0 hours, 1 hours, 2 hours, 3 hours, and 4 hours (6).

2 LPS brought on fever in Wistar rats

The animals have been randomly divided into 5 groups (n = 6). Animals in Groups 1, obtained the automobile pyrogen-unfastened water and 2d received10 mg/kg frame weight of indomethacin. And institution 0.33 and fourth obtained cinnamon oil 25 and 50 miligram in line with kilogram orally. Before beginning the test all institution animals have been injected intraperitoneally with LPS (500 mg/kg) dissolved in pyrogen-unfastened sterile regular saline solution. Body temperature (Tb) turned into taken each h for five h after LPS administration. These experiments have been done throughout the mild segment of the circadian cycle. Temperature turned into measured with the aid of using lightly putting a small thermoprobe approximately four cm into the rectum till a beep sound heard. During the temperature measurements, every animal turned into held lightly and done at the least two times 2 days previous to the test to keep away from adjustments in rectal temperature secondary to handling. Only animals with strong frame temperature and within the variety 32.four– 37.zero _C have been used within side the research of the have an effect on of the drug at the Tb of the animals upon induction of pyrexia with LPS (7).

Statistical observation

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The statistical investigation was finished using Graphpad 5.0 programming. Information was noted as mean ± S.E.M. The measurable outcome of variety between bunches was finished up by the examination of various two way’s (ANOVA) followed by means of Bonferroni’s posttests. Contrasts of P<0.001 were analyzing statically demonstrative

IV. RESULTS AND DISCUSSION

The impact of cinnamon oil on the rectal degree of heat in rodents is depicted in (Table-1). The sub-cutaneous infusion of brewer's yeast’s suspensions uniquely expanded the rectal degree of heat after 18h of the organization. prevention with cinnamon oil 25mg or 50 milligram/kilogram diminished the colonic heat of the rat in portion subordinate kind. It was discovered that the cinnamon oil 25 Milligram/ kilogram created significant bringing down the internal heat level at 4 hours following its organization (96.42 ± 0.009). This impact was maximal at a portion of 50 Milligram/kilogram at which created significant bringing down the internal heat level (P<0.01) as long as 4 hours after its organization (95.40 ± 0.017). The antipyretic activity began as ahead of schedule and 1 h and the impact were kept up for 4h, after its organization. Standard medication paracetamol 100mg/kg and test tranquilize (cinnamon oil) were essentially lessened the yeast-raised rectal temperature, at second, third, and fourth hour contrasted with the benchmark group.

Table 1- Antipyretic Effect of Cinnamon oil on Brewer’s Yeast Influenced warmth in Rat

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Dose(milligram/kilogram)</th>
<th>Basel heat (.°F)</th>
<th>0 hour’s (after18hr)</th>
<th>one hour’s</th>
<th>Two-hour’s</th>
<th>Three hour’s</th>
<th>Four hour’s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group</td>
<td>-</td>
<td>95.57 ± 1.223</td>
<td>101.27 ± 0.414</td>
<td>101.12±0.007</td>
<td>101.08±0.058</td>
<td>101.19±0.009</td>
<td>101.08±0.007</td>
</tr>
<tr>
<td>Paracetamol Standard group</td>
<td>100mg/kg</td>
<td>95.42±0.007</td>
<td>100.17±0.008</td>
<td>98.24±0.006 ***</td>
<td>97.21±0.008 **</td>
<td>96.50±0.008 ***</td>
<td>95.49±0.403 ***</td>
</tr>
<tr>
<td>Cinnamon oil Test group-I</td>
<td>25mg/kg</td>
<td>96.12 ± 0.008</td>
<td>101.08 ± 0.006s</td>
<td>99.02.12 ± 0.005***</td>
<td>97.51 ± 0.017 **</td>
<td>96.78 ± 0.009 ***</td>
<td>96.42 ± 0.009 **</td>
</tr>
<tr>
<td>Cinnamon oil Test group-II</td>
<td>50mg/kg</td>
<td>96.22±0.409</td>
<td>100.47 ± 0.012</td>
<td>99.43±0.005***</td>
<td>97.48 ± 0.407 **</td>
<td>96.20 ± 0.005 ***</td>
<td>95.40 ± 0.017 **</td>
</tr>
</tbody>
</table>

Figure-3 Line Graph Displaying the Effect of Cinnamon oil 25mg/kg and 50mg/kg Against Brewer’s Yeast’s Instigated
Warmth

Table 2- Anti-pyretic consequence of Cinnamon oil Lipopolysaccharide produced warmth in Rat

<table>
<thead>
<tr>
<th>Treatment Dose(milligram/kilogram)</th>
<th>Basel Heat.(°F)</th>
<th>Colonic heat (°F)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 hour (after 5hr)</td>
<td>1 hour</td>
</tr>
<tr>
<td>Control group</td>
<td>-</td>
<td>95.59 ± 1.221</td>
</tr>
<tr>
<td>Indomethacin standard group</td>
<td>10mg/kg</td>
<td>95.40±0.005</td>
</tr>
<tr>
<td>Cinnamon oil Test group-I</td>
<td>25mg/kg</td>
<td>96.11 ± 0.006</td>
</tr>
<tr>
<td>Cinnamon oil Test group-II</td>
<td>50mg/kg</td>
<td>96.22±0.401</td>
</tr>
</tbody>
</table>

The impact of cinnamon oil on the rectal degree of heat in rodents is depicted in (Table-5). intraperitoneally with LPS (500 mg/kg) suspension uniquely expanded the rectal degree of heat after 5 hours of the organization. prevention with cinnamon oil 25mg or 50 milligram/kilogram diminished the colonic degree of the rat in portion subordinate kind. It was discovered that the cinnamon oil 25 Milligram/ kilogram created significant bringing down the internal heat level at 4 hours following its organization (96.45 ± 0.009). This impact was maximal at a portion of 50 Milligram/ kilogram at which created significant bringing down the internal heat level (P<0.01) as long as 4 hours after its organization (95.40 ± 0.000). The antipyretic activity began as ahead of schedule and 1 h and the
impact were kept up for 4h, after its organization. Standard medication paracetamol 100mg/kg and test tranquilize (cinnamon oil) were essentially lessened the yeast-raised rectal temperature, at second, third, and fourth hour contrasted with the benchmark group.

V. CONCLUSION

Pyrexia and agony are considered as an auxiliary effect of contamination tissue harm, aggravation, join dismissal, threat, or another infected state. It is the body’s characteristic barrier to make a situation where irreversible operators or harmed tissue can’t endure. A large portion of the Non steroidal anti-inflammatory medication hinders Cyclo-oxygenase-2 articulation to lessen the raised body’s degree of heat by restraining PGE2 biosynthesis. Besides, these manufactured operators irreversibly hinder Cyco-oxygenase type-2 Basic Cyclo-oxygenase-2 blockers, on either hand, have a lesser preference and having low reflexes, but they are dangerous to liver tissues, nephron’s, the cortex of the mind, and cardiacmuscles [8] Since antipyretic potentiality is frequently reported as a feature of drugs or compounds, which have an inhibitory action on prostaglandin biosynthesis, the yeast produced hyperpyrexia in a rat model was applied to explore the antipyretic activity of the cinnamon,[9] Yeast instigated fever is called pathogenic fever which is due to the formation of Prostaglandins-E2, who enhance the heat of the thermo-regulatory centre. The current pharmacological research was exploring the antipyretic capability of cinnamon oil (25 mg/kg and 50mg/kg).Cinnamon oil shows a significant anti-Pyretic effect in Wistar rodents after oral administration. So we can use cinnamon oil for the management of pyrexia [10].

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Author contribution

All author participated Equally.

Conflict of interest None

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REFERENCE