Pharmacological Study

Toxicological evaluation of Panchakola Avaleha, an Ayurvedic classical formulation, in albino rats

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Abstract

The present study was carried out to assess the safety of standardized Panchakola Avaleha on albino rats (Wistar strain). Animals were administered three doses of Panchakola Avaleha by oral routes, viz. higher (500 mg/kg/day), middle (250 mg/kg/day), and therapeutic dose (50 mg/kg/day) for 28 consecutive days. Effects of the test drug on hematological, biochemical, and histopathologic parameters were evaluated. This study revealed normal behavior, no mortality, and no significant changes in hematological, biochemical, and histopathological examinations.

Key words: Panchakola Avaleha, toxicity study, Wistar rats

Introduction

Panchakola is one of the most popular formulations in Ayurveda, which is being used as a general health tonic. It is used as antipyretic, analgesic, anti-inflammatory, appetizer, digestive, and carminative.[1] It is highly effective particularly in Vatasleshmika Jvara.[2] It is available in different pharmaceutical forms like Churna (powder), Paka or Avaleha (confection), and Ghrita (medicated fatty preparation). Panchakola Avaleha rejuvenates the female reproductive system, particularly in periparum stage and helps in early involution of uterus. This formulation also helps to improve the quality and quantity of breast milk.[3] A recent clinical study showed that Panchakola Siddha Yavagu is effective as an appetizer.[4] This formulation is also being used for centuries in different conditions of Sandhigata Vata (Arthritis and rheumatism) as a supportive measure. In fact, the toxicological study of Panchakola Avaleha has not been done and the present study is being carried out for toxicological evaluation. This article presents the results of a 28-day repeated dose oral toxicity study conducted to ensure the safety of the drug.

Materials and Methods

This study was conducted at National Research Institute of Ayurvedic Drug Development, Kolkata, a peripheral center under Central Council for Research in Ayurvedic Sciences, Department of AYUSH, Ministry of Health and Family Welfare, Government of India.

Test animals and housing

The Institutional Animal Ethics Committee (IAEC) approved the experimental protocol (138/ARI/2000-2002) dated 29.12.2004 and Registration No. 694/a/CPCSEA/03 dated 01.10.2002. The present study was conducted on inbred albino rats (Wistar strain) of both sexes, which were procured from NIN, Hyderabad.

Wistar rats of 6-7 weeks of age, weighing 140-250 g (20 males + 20 females = 40) were selected based on the body weight and distributed into four groups (5 males and 5 females in each group). Animals were acclimatized for 7 days and health examination was performed during acclimatization period. Rats were housed individually in polycarbonate cages, were fed animal pellet diet and mineral water ad libitum during the entire study period.

The temperature was maintained at 26 ± 3°C and relative humidity at 60-70%, and illumination was controlled to give approximately a sequence of 12 h light and 12 h dark.

Test drug

The test drug, i.e., Panchakola Avaleha is composed of five major indigenous plant materials, viz. Pippali (Piper longum L.), Pippalimula (Piper longum L.), Chavya (Piper chaba Trel. and Yunck.), Chitraka (Plumbago zeylanica L.), and Shunthi (Zingiber officinale Roscoe), and Guda (jaggery), Sarkara (sugar) and Ghrita (ghee) were used for preparation of Avaleha following the method of preparation of Ayurvedic practice and principles[5] [Table 1].
Experimental design
Twenty-eight days repeated dose oral toxicity study was carried out on 40 rats (both sexes) and the rats were divided into four groups consisting of 5 males and 5 females in each. The rats received daily vehicle control (honey:Deionized water 2:3 proportion) and 500, 250, and 50 mg/kg/day Panchakola Avaleha (dissolved in honey:Deionized water 2:3 proportion) by oral gavage, once daily for 28 consecutive days. Both the test and control groups received the same volume of drug or vehicle as per body weight. Animals were checked for mortality, and general clinical observations, viz. salivation, activity, irritability, fecal pellet condition, diarrhoea, eye ball movement, and external appearance, were made daily in the morning and evening. The body weight was taken before the start of the treatment in vehicle control as well as in treated groups. The weekly feed consumption and water consumption of rats were recorded by measuring the difference between the feed and water offered and feed and water left over the subsequent week. The weekly feed and water consumed per cage was calculated and presented. Blood sampling was performed on day 29, prior to euthanasia. Animals were fasted overnight, and blood samples were drawn from the retro-orbital plexus under diethylether-induced anesthesia. For hematological analysis, blood was collected in a silica-coated vial with anticoagulant containing Ethylene Di Amine Tetra Acetic acid (EDTA) to measure hemoglobin (HB%), Total Erythrocyte Count (TEC), and Total Leukocyte Count (TLC). Blood was collected in a centrifuge tube without anticoagulant for collection of serum for clinical chemistry parameters including serum total bilirubin,[8] total protein, serum albumin,[7] Serum Glutamic Pyruvic Transaminase (SGPT), Serum Glutamic Oxaloacetic Transaminase (SGOT),[8] serum alkaline phosphatase,[9] and serum creatinine.[7] Necropsy was performed on day 29 of the study. Organs were observed, collected, weighed, and preserved in 10% neutral buffered formalin, viz. kidney, adrenal, heart, spleen, and liver. Tissues were then trimmed and dehydrated in ascending grades of alcohol. Finally the tissues were embedded in melted paraffin and blocks were prepared. The tissue sections were cut to 3-5 μm in microtome and stained with hematoxylin and eosin.[10] Finally, the sections were mounted with Distyrene Plastizeric and Xylene (DPX) and examined under microscope. Histopathologic examination was performed on all collected tissues from males and females of the control and 500 mg/kg/day group.

Statistical analysis
All the data were analyzed by Student’s t-test followed by analysis of variance (ANOVA).

Observations and Results
No mortality and treatment-related clinical signs were observed throughout the study period. There was no significant change in treated animals at 2nd and 4th weeks, when compared with the initial values of the respective groups. However, the mean values of treated animals were higher when compared with controlled group and these changes were attributed to sporadic deviation within the groups and had no biological significance, hence not considered as drug-related adverse effect [Table 2]. All animals from Panchakola Avaleha treated groups consumed similar amount of food, compared with the corresponding control group. The observations of feed and water intake changes were not dose dependent and not observed consistently. Hence, it was not considered as a treatment-related effect. In case of treated animals in 50 mg/kg/day group, statistically significant increase of hemoglobin percentage on 25th day was observed; however, the value was within normal range [Table 3]. There were no significant changes in liver function tests, i.e., total serum bilirubin, SCOT, SGPT, serum albumin, total protein, alkaline phosphatase, and serum creatinine, on 28th day of study when compared with their values in the respective groups and control group, except for the levels of albumin and total protein in 500 mg/kg/day group and the level of total protein in 250 mg/kg/day group [Table 4]. There were also no significant changes in weights of vital organs, viz. kidney, adrenal, heart, spleen, and liver, on 28th day of study [Table 5]. Gross necropsy examination revealed no treatment-related lesions. There was significant increase in weight of liver and spleen in 500 mg/kg/day group, but in other groups no such change was observed [Table 5]. Hence, it is not considered as a treatment-related effect. However, histopathology evaluation of any of the organ did not reveal any treatment-related or dose-dependent changes [Figures 1-10].

Discussion
Among the variety of Ayurvedic medicines, drugs are being used to treat various diseases according to Ayurvedic pharmacology. Since this traditional treatment is based on extensive knowledge gathered from applications of natural resources to humans, people have usually assumed that the treatment is safe.

### Table 1: Composition of Panchakola Avaleha in 10 g

| Name of the ingredients | Botanical name | Part of use | Amount (mg) |
|-------------------------|----------------|-------------|-------------|
| Pippali                 | Piper longum   | Fruit       | 435.0       |
| Pippalimula             | Piper longum   | Root and stem | 435.0    |
| Chavya                  | Piper chaba    | Stem        | 435.0       |
| Chitraka                | Plumbago zeylanica | Root  | 435.0       |
| Shunthi                 | Zinger officinale | Rhizome   | 435.0       |

Guda (jaggery) – – 2250.0
Sarkara (sugar) – – 5050.0
Ghrita (ghee) – – 525.0

Preservatives such as sodium methyl paraben (0.15%), propyl paraben (0.05%), sodium benzoate (0.3%), and antioxidant (0.1%) were used in the formulation.

### Table 2: Effect of Panchakola Avaleha on body weight (g) measured weekly in rats at different doses for 28 days (n=10)

| Body weight (vehicle) group | Test drug groups-PKA (mg/kg) |
|-----------------------------|------------------------------|
|                             | 500  | 250  | 50   |
| Initial wt.                 | 170.0±05.16  | 183.0±11.75  | 181.0±09.48  | 172.0±07.72  |
| 1st week                    | 177.0±09.20  | 195.0±11.18  | 180.0±10.86  | 179.0±09.00  |
| 2nd week                    | 186.0±09.45  | 203.0±14.92  | 197.0±12.75  | 190.0±12.21  |
| 3rd week                    | 187.0±09.19  | 208.0±26.03  | 198.0±24.02  | 194.0±21.64  |
| 4th week                    | 198.0±04.91  | 226.0±28.99  | 199.6±25.48  | 200.0±29.23  |

Data represented as mean±SEM; PKA - Panchakola Avaleha
Table 3: Investigation of hematology of rat blood treated with *Panchakola Avaleha* on 28th day (n=10)

| Hematological parameters | Control (vehicle) group | Test drug groups-PKA (mg/kg) |
|--------------------------|-------------------------|-------------------------------|
|                          |                        | 500                           |
|                          |                        | 250                           |
|                          |                        | 50                            |
| Hb (g% /dl)              | 16.20±1.43             | 18.62±2.19                    |
| WBC (10³/mm³)            | 06.62±0.18             | 06.42±0.32                    |
| RBC (10⁹/mm³)            | 06.04±0.34             | 05.48±0.64                    |
| Leukocyte                | 61.00±1.67             | 66.40±1.75                    |
| Neutrophil               | 32.60±1.43             | 28.40±1.50                    |
| Monocyte                 | 01.00±0.31             | 00.60±0.24                    |
| Eosinophil               | 05.00±0.31             | 04.40±0.40                    |
| Basophil                 | 00.42±0.24             | 00.20±0.20                    |

Data are represented as mean±SEM; PKA - *Panchakola Avaleha*; *P*<0.05 (control vs. treated)

Table 4: Investigations of blood biochemistry of rat treated with *Panchakola Avaleha* on 28th day (n=10)

| Bio-chemical parameters | Control (vehicle) group | Test drug groups-PKA (mg/kg) |
|-------------------------|-------------------------|-------------------------------|
|                         |                        | 500                           |
|                         |                        | 250                           |
|                         |                        | 50                            |
| Total bilirubin (mg/dl) | 1.08±0.35              | 1.17±0.30                     |
| SGOT (U/L)              | 144.80±2.79            | 147.60±17.58                  |
| SGPT (U/L)              | 46.40±4.23             | 52.40±6.68                    |
| Albumin (g/dl)          | 2.90±0.44              | 4.30±0.24*                    |
| Serum creatinine (mg/dl)| 0.67±0.09              | 0.58±0.05                     |
| Alkaline phosphatase (U/L) | 189.80±31.44 | 260.60±55.18                  |

Data are represented as mean±SEM; PKA - *Panchakola Avaleha*; *P*<0.05 (control vs. treated)

Table 5: The weight of vital organs (g) treated with *Panchakola Avaleha* on 28th day (n=10)

| Vital organs (vehicle) group | Test drug groups-PKA (mg/kg) |
|-----------------------------|-------------------------------|
|                             | 500                           |
|                             | 250                           |
|                             | 50                            |
| Heart                      | 0.66±0.03                     |
| Kidney                     | 1.32±0.05                     |
| Adrenal                    | 0.056±0.006                   |
| Spleen                     | 0.42±0.01                     |
| Liver                      | 6.53±0.33                     |

Data are represented as mean±SEM; PKA - *Panchakola Avaleha*; *P*<0.05 (control vs. treated)

Conclusion

It is revealed from the study that there was no mortality of the experimental animals. There were no morphological, haematological and biochemical changes in the treated animals when compared with control animals. It may be concluded that *Panchakola Avaleha* is safe in animal models.

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हिन्दी सारांश
शास्त्रीय आयुर्वेदिक औषध योग-पंचकोल अवलेह के विषाक्तता प्रभाव का प्रायोगिक अध्ययन

राजेन्द्र कुमार सिंह, रिता बनर्जी, सचिदानंद उपाध्याय, अचिन्त्य मित्र, जयराम हाजरा

मानकीकृत पंचकोल अवलेह के विषाक्तता प्रभाव का सुरक्षात्मक मूल्यांकन चूक़ों पर किया गया। मुख्य रूप से पंचकोल अवलेह औषध की तीन प्रकार की मात्रा का चूकों पर अध्ययन - उब (५०० मि.ग्रा./कि.ग्रा./दिन), मध्य (२५० मि.ग्रा./कि.ग्रा./दिन) और निम्न (५० मि.ग्रा./कि.ग्रा./दिन) लगातार २५ दिनों तक दिया गया। इस औषध का प्रभाव, रक्त एवं विभिन्न अंगों पर प्रभाव का हिस्टोपैथोलॉजी अध्ययन किया गया। इस अध्ययन में सभी प्राणियों में सामान्य व्यवहार, मृत्यु दर एवं उसका रक्त और विभिन्न अंगों पर कोई विषाक्त प्रभाव नहीं पाया गया।