**In vitro Evaluation of Anthelmintic Activity of Gymnema sylvestre Plant**

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**ABSTRACT**

The evaluation of anthelmintic activity of *Gymnema sylvestre* was performed by bioassay method where hot and cold hydroalcoholic extracts were used against earthworm (*Pheretima posthuma*). The anthelmintic assay was carried out as the 50ml formulations containing five different concentrations of each cold and hot hydroalcoholic extracts (25, 50, 100, 250 and 500mg/ml in distilled water) were prepared and six worms (same type) were placed in them. Time for paralysis and death time was noted. Albendazole (20mg/ml) was used as a reference standard, while normal saline as the control.

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**INTRODUCTION**

Helminths are parasitic worms, they are most common infectious agents of humans in developing countries and produce a global burden of disease. Helminths have plagued humans since before the era of our earliest recorded history.¹ There are two major phyla of helminths. The nematodes (roundworms) include the major intestinal worms, also known as soil transmitted helminths. Whereas platyhelminths (flatworms) include flukes also known as trematodes and tapeworms. It is estimated that approximately one-third of almost three billion people that live on less than two US dollars per day in developing regions of Sub-saharan Africa, Asia, and America are affected with one or more helminth.² The high medical, educational and economic burden of helmint infections provides an important rationale for launching a global assault on parasitic worms.³ However, the tools we currently have for controlling worm infections are limited, of the 1,556 new chemical entities currently have for controlling worm infections evaluated in the development and discovery of new anthelmintic produced toxicity in humans. Hence the development and discovery of new anthelmintic are being derived through plants. Herbal medicine is still the main source of medicine and about 70-80% of the whole population, mainly in developing countries for primary healthcare because of better cultural acceptability, better compatibility, with the human body and fewer side effects. *Gymnema sylvestre* also known as madhunashini (Sanskrit) and cherukurinja (tamil) belonging to the family of Asclepiadaceae was a vulnerable species is a slow growing, perennial, medicinal woody climber found in central and peninsular India. It is a potent anti-diabetic plant and used in folk, ayurvedic and homeopathic systems of medicine.⁴ In addition, it poses wound healing,⁵ anti-inflammatory,⁶ anti-obesity⁷ treatment of snake bite⁸ and anthelmintic properties. Flavonoids, tannins and saponins are the chief constituents of *Gymnema sylvestre*.⁹,¹⁰,¹¹ *Gymnema sylvestre* is used in ayurvedic medicine as anthelmintic but it is not clinically proved. The active principles are flavonoids which show selective anthelmintic activity.

**MATERIALS AND METHODS**

**Drugs and chemicals**

Albendazole, normal saline (Abaris healthcare private limited) and ethanol.

**Plant collection and authentication**

The leaves of *Gymnema sylvestre* belonging to the family of Asclepiadaceae were collected from the herbal garden of Sri Ramachandra Institute of Higher Education and Research, Porur, Chennai. The plant was authenticated Dr. P. Jayaraman (Director, Plant Anatomy Research Centre, Tambaram).

**Plant extraction**

Leaves were dried at room temperature (25 to 35 ºc) and powdered with the help of a electrical grinder. The fine material was subjected to extraction, successfully by hot and cold hydroalcoholic mixture in the ratio of (water60:40 ethanol) as solvent. The extract was allowed to dry at 100 ºC in water bath. The percentage yield of the different successive extracts was 30 and 40% respectively.

**Worms collection**

Indian earthworms *Pheretima posthuma* of nearly equal size (8 to 10 cm) were collected from the waterlogged areas from herbal garden of SRI RAMACHANDRA INSTITUTE OF HIGHER EDUCATION AND RESEARCH.

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Preparation of test sample

Samples for in vitro study were prepared by dissolving and suspending (0.12,0.25,0.5,1.25 and 2.5g) of each hot and cold hydroalcoholic extract in 50ml of distilled water at different concentrations ranging from 25, 50, 100, 250 and 500 mg/ml.

Anthelmintic assay

It was carried out using adult earthworm (Pheretima posthuma) owing to its anatomical and physiological resemblance with the intestinal roundworm parasites of human beings for preliminary evaluation of anthelmintic activity.13 The 50ml formulations containing five different concentrations of each hot and cold hydroalcoholic extracts (25,50,100,250 and 500mg/ml in distilled water) were prepared. Pheretima posthuma was placed in petri dish containing 10ml of the extract.14 Each petri dish was placed with six worms and observed for paralysis and death. The mean time for paralysis was noted when no movement of any sort could be observed, except when the worm is shaken vigorously. The time of death of worms was recorded after ascertaining that the worms neither moved when shaken vigorously nor when dipped in normal saline followed with fading away of their body colour and the results were expressed in comparison to the standard drug Albendazole (20 mg/ml).

RESULTS AND DISCUSSION

From the Table 1, it was evident that hot and cold hydroalcoholic extracts of Gymnema sylvestre exhibited anthelmintic activity in dose dependent manner giving the shortest time of paralysis and death with 500 mg/ml concentration. The hot extract caused paralysis in 15 minutes and time of death 23 minutes, while cold extract revealed paralysis of 24 minutes and time of death 35 minutes against the earthworm Pheretima posthuma.

Chemically flavonoids are polyphenolic compounds and they interfere with the energy generation by uncoupling the oxidative phosphorylation which interfere with the glycoprotein of cell surface leads to parasite death.15 Another possible anthelmintic effect of tannins is that they bind to free proteins in the gastrointestinal tract of host animal or glycoprotein on the cuticle of the parasite and may cause death.16

HOT EXTRACT

From the Figures 1-4, Albendazole standard (20 mg/ml) showed paralysis time in 42 ± 0.2 min and death time in 54.15 ± 0.3 min. In hot extract 25mg showed paralysis time in 53 ± 0.5 min and death time in 61.02 ± 0.4 min, 50mg showed paralysis time in 49.2 ± 0.2 min and death time in 57.18 ± 0.6 min, 100mg showed paralysis time in 37.23 ± 0.4 min and death time in 43.13 ± 0.3 min, 250mg showed paralysis time in 21.07 ± 0.6 min and death time in 29.11 ± 0.4 min and 500mg showed paralysis time in 15.17 ± 0.4 min and death time in 23.02 ± 0.6 min.

COLD EXTRACT

From Figure 5, In cold extract 25mg showed paralysis time in 55.11 ± 0.2 min and death time in 64.03 ± 0.3 min, 50mg showed paralysis time in 51.33 ± 0.1 min and death time in 59.12 ± 0.7 min, 100mg showed paralysis time in 45.19 ± 0.3 min and death time in 55.05 ± 0.4 min, 250mg showed paralysis time in 36.16 ± 0.8 min and death time in 49.06 ± 0.6 min and 500mg showed paralysis time in 24.17 ± 0.3 min and death time in 35.11 ± 0.2 min.

Table 1: In vitro evaluation of anthelmintic activity of Gymnema sylvestre.

| Group | Solution   | Concentration in Mg/Ml | Time taken for paralysis (Mins) | Time taken for death (Mins) | Time between paralysis and death (Mins) |
|-------|------------|------------------------|---------------------------------|-----------------------------|----------------------------------------|
| 1.    | Control    | -                      | -                               | -                           | -                                      |
| 2.    | albendazole| 20                     | 42.2 ± 0.2                      | 54.15 ± 0.3                 | 11.95 ± 0.1                            |
| 3.    | Hot extract| 25                     | 53.05 ± 0.3                     | 61.02 ± 0.4                 | 7.97 ± 0.1                             |
|       |            | 50                     | 49.12 ± 0.2                     | 57.18 ± 0.6                 | 8.06 ± 0.4                             |
|       |            | 100                    | 37.23 ± 0.4                     | 43.13 ± 0.3                 | 5.9 ± 0.1                              |
|       |            | 250                    | 21.07 ± 0.6                     | 29.11 ± 0.4                 | 8.04 ± 0.2                             |
|       |            | 500                    | 15.17 ± 0.4                     | 23.02 ± 0.6                 | 7.85 ± 0.2                             |
| 4.    | Cold extract| 25                    | 55.11 ± 0.2                     | 64.03 ± 0.3                 | 8.92 ± 0.3                             |
|       |            | 50                     | 51.33 ± 0.1                     | 59.12 ± 0.7                 | 7.79 ± 0.6                             |
|       |            | 100                    | 45.19 ± 0.3                     | 55.05 ± 0.4                 | 9.86 ± 0.1                             |
|       |            | 250                    | 36.16 ± 0.8                     | 49.06 ± 0.6                 | 12.9 ± 0.2                             |
|       |            | 500                    | 24.17 ± 0.3                     | 35.11 ± 0.2                 | 10.94 ± 0.1                            |
Figure 2: *In vitro* evaluation of anthelmintic activity of *Gymnema sylvestre* using hot extract.

Figure 3: *In vitro* evaluation of anthelmintic activity of *Gymnema sylvestre* using cold extract.

Figure 4: Administration of hot extract in various concentrations.
Figure 5: Administration of cold extract in various concentrations.

REPORT

The graph of hot extract and cold extract expresses the paralysis time and death time at various concentrations of the extract which exhibited anthelmintic activity in dose dependent manner giving the shortest time of paralysis and death with 500 mg/ml concentration. The graph helped to identify the concentration at which the paralysis and death occurs in the shortest duration of time. In hot extract the hot extract (Figure 2) caused paralysis in 15 minutes and time of death 23 minutes, while cold extract (Figure 3) revealed paralysis of 24 minutes and time of death 35 minutes against the earthworm Pheretima posthuma.

CONCLUSION

From the results, it was concluded that both hot and cold hydroalcoholic extracts of Gymnema sylvestre have significant anthelmintic activity, but hot hydroalcoholic extract shown most significant anthelmintic activity when compared to cold hydroalcoholic extract. From the results, Gymnema sylvestre has an anthelmintic activity have been confirmed as it displayed activity against the worm used in the present study.

FUTURE SCOPE

Further studies have to be done to isolate the active principles responsible for the activities. In addition to that we are going to perform comparative studies on various species of Gymnema sylvestre have significant anthelmintic activity, but hot hydroalcoholic extract shown most significant anthelmintic activity when compared to cold hydroalcoholic extract. From the results, Gymnema sylvestre has an anthelmintic activity have been confirmed as it displayed activity against the worm used in the present study.

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**GRAPHICAL ABSTRACT**

**ANTHELMINTIC ACTIVITY**

- Preparation of Extract
  - Leaves of *Gymnema sylvestre*
  - Hot extract (Percolation)
  - Cold extract (Maceration)
  - % yield of hot extract = 30% W/W
  - % yield of Cold extract = 40% W/W

**ANALYSIS**

- Hot extract
- Cold extract

**Death and paralysis time calculated using Hot and cold extract against Pheretima posthuma**

- Hot hydroalcoholic extract shown most significant anthelmintic activity when compared to Cold hydroalcoholic extract.

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My career started in professional service as Lecturer and continued as Professor & Principal in reputed institutions. In nutshell, has 27 years of teaching experience and 20 years of research experience, Ph.D awarded from Sri Ramachandra University, Chennai, As principal investigator /Co investigator obtained grants from Govt funded bodies. As a credit, 24 international Publications, chapter contributions and presented research output in 85 International & National conferences. As a member various professional bodies Member, Board member of various University, National Advisory Committee member, Society for Ethnopharmacology. Evaluator for PhD thesis for various Universities. As a research potential has been witnessed by various awards received from Society For Ethnopharmacology, Universities and Govt bodies and research funds from Dept. of AYUSH and National Medicinal Plant Board. Chaired many International and National conferences and served as resource person too. Organized good number of national & International conferences in the field of Pharmacognosy.

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Currently Professor in Pharmacognosy, with teaching experience of 29 years. As principal investigator /Co investigator obtained grants from DST, DBT NER, AYUSH, ILSRB-DRDO and SPARC worth Rs. 8.5 crores. About 95 publications with impact factor of 38 in Scopus, filed 6 patents, 2 published and 1 granted. As member of the curriculum committee of PCI, designed curriculum for M.Pharm (Pharmacognosy). As a team edited text book of Pharmacognosy authored by Shah and Quadry. Awarded Young Scientist in 2007, by DST, New Delhi. As expert in Siddha Pharmacopoeial Committee, CCRS, Chennai contributed to the development of Volume II of Siddha Pharmacopoeia.
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