Evaluation of Comparative Dermal Toxicity and Efficacy of Aargwadhadi Oil and Aargwadhadi Ointment

Premadevi Kalmegh a#, Bharat Rathi b*†, Renu Rathi c† and Poonam Madan d‡

a Department of Rasashastra & Bhaishajya Kalpana, Rajendra Gode Ayurveda College Hospital and Research Centre, Amravati, MGACHRC, DMIMS (DU), India.
b Department of Rasashastra & Bhaishajya Kalpana, Mahatma Gandhi Ayurveda College Hospital & Research Centre, Datta Meghe Institute of Medical Sciences (DU), Sawangi (M) Wardha, India.
c Department of Kaumarbhritya, Mahatma Gandhi Ayurveda College Hospital & Research Centre, Datta Meghe Institute of Medical Sciences (DU), Sawangi (M) Wardha, India.
d Department of Rasashastra & Bhaishajya Kalpana, Datta Meghe Ayurveda College Hospital & Research Centre, Wanadongari, Nagpur, India.

Authors’ contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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ABSTRACT

Introduction: Skin diseases are among the most common of all health illness in recent years, In Ayurveda, management of skin diseases includes internal and external administration. External administration includes various topical applications However, herbal medicines are not completely safe from adverse effects and develop irritation, rashes, redness and burning sensation on the skin as observed in recent researches. “Hence there is a requirement of a safe drug which should be economical & affordable for all. Hence present study is undertaken to study dermal toxicity profile Aargwadhadi oil and Aargwadhadi ointment in an experimental animal and make available safe and efficient drug to the human being.

Aim and Objective: Pharmaceutical development, standardization and evaluation of acute, sub-
Materials and Methods: Aargwadhadi oil and Aargwadhadi Ointment will be prepared as per classical reference and it will be converted into ointment form. Analytical study for standardization of Aargwadhadi oil and Aargwadhadi ointment will be done. Evaluation of Acute and sub-acute dermal toxicity study in experimental animals of both dosage forms as well, efficacy study of Aargwadhadi oil and Aargwadhadi ointment on animal model of vitiligo will be done.

Observations and Results: Observation will be done on the basis of assessment criteria evaluation of control group and experimental group will be noted. Results will be drawn on the basis of observations and applying suitable tests. It will be noted and presented in form of table, charts, graphs etc.

Conclusion: Conclusion of the study will be drawn accordingly from the recorded observations, analysis of data.

Keywords: Aargwadhadi oil; Aargwadhadi ointment; acute & sub-acute dermal toxicity; vitiligo.

1. INTRODUCTION

Ayurveda is a complete and holistic traditional health-care system which have both preventive and therapeutic aspects. Ayurveda has eight specialized branches called as Ash tang Ayurveda. The enduring fundamentals of Ayurveda, which were said by various Acahryas are valid and still applicable because of their scientific background. In today’s technology and scientific era these fundamentals must be subjected to scientific research not only to prove it but also to add something new to the existing knowledge and further in the betterment of health [1].

Acharya Charka has mentioned that any materials in the universe or even a poison can be used as a medicine when it is used properly and on the contrary improperly used medicine can act as a poison [2]. Traditionally medicinal plants have been used for many years as topical and internal preparation in the treatment of fungal and bacterial diseases. Globally, skin conditions are the fourth leading cause of nonfatal disease burden at the global level [4-5]. Various studies have shown that skin diseases can have a major impact on the quality of life of those affected [6]. Some skin diseases, such as skin cancer and infections are potentially life-threatening [7].

Safety is the most important consideration before administration of any herbal product in human being. In Ayurveda, management of skin diseases includes internal and external administration. External administration includes various Lepas (Paste), medicated oil, poultice, face pack, hair pack etc which are advised to apply on affected skin for varied duration [8-9]. However, herbal medicines are not completely safe from adverse effects and develop irritation, rashes, redness and burning sensation on the skin as observed in recent researches [10-12].

Though the modern medicine is having powerful antibiotics, antifungals, antihistamines, steroids etc., but better management could not be searched out till today. Moreover, topical medicines are expensive and sometimes poor people cannot afford, hence there is a need to search safe drug which is economical and affordable to the society. The adverse reaction of modern cosmetics and higher cost of therapy are also made the society to approach ancient system of Ayurveda.

Hence present study is undertaken to study dermal toxicity profile Aargwadhadi Oil and Aargwadhadi ointment in an experimental animal and make available safe drug to the human being.
1.1 Aims and Objective of Study

1.1.1 Aim of study
Evaluation of Comparative Dermal Toxicity and Efficacy of Aargwadhadi Oil and Aargwadhadi Ointment.

1.1.2 Objectives of the study
1. To prepare Aargwadhadi Oil and Aargwadhadi Ointment by standard operating procedure
2. To evaluate acute dermal toxicity of Aargwadhadi Oil and Aargwadhadi Ointment in experimental animal.
3. To evaluate sub-acute dermal toxicity of Aargwadhadi Oil and Aargwadhadi Ointment in experimental animal.
4. To evaluate efficacy of Aargwadhadi Oil and Aargwadhadi ointment in vitiligo animal model.
5. To compare safety and efficacy of Aargwadhadi Oil and ointment

2. MATERIALS AND METHODS
Present work will be conducted under following headings.

2.1 Pharmaceutical Study
In this study Aargwadhadi Oil and Aargwadhadi Ointment will be prepared in three batches as per classical reference to establish pharmaceutical standardization. This pharmaceutical study will be done according to following steps.

- Procurement of raw drug – all raw drugs will be collected / procured from the field and authentic reliable sources.
- Authentication of raw material – all raw materials will be verified and authenticated by department of Dravyaguna of the institute. Raw drug will be standardized as per A.P.I. Table 1 show contents of Aargwadhadi Oil and Aargwadhadi ointments and their quantity. Below is the flow diagram (Fig. 1) of method of preparation of Aargwadhadi Oil and Aargwadhadi ointment.

2.2 Analytical Study [14-17]

2.2.1 Analytical study of Aargwadhai Oil
i. Organoleptic characters
- Sparsha (touch), Rupa (appearance), Gandh (odour) will be assessed

ii. Physicochemical parameters
1. Specific Gravity
2. Refractive index at 25°C
3. Viscosity
4. Acid value
5. Saponification value
6. Iodine value
7. Peroxide value
8. Unsaponifiable matter
9. HPTLC or GC-MS (Quantitative)

2.3 Preparation of Aargwadhadi Ointment
Aargwadhadi Oil will be made into ointment by following SOP. Table 2. shows ingredients and quantity and below is the flow diagram (Fig. 2) of method of preparation of Aargwadhadi ointment.

Table 1. Ingredients used for Aargwadhadi Oil [13]

<table>
<thead>
<tr>
<th>Sr no</th>
<th>Drug Name</th>
<th>Part used</th>
<th>Proportion</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Aargwadh (Cassia fistula)</td>
<td>Fruit pulp</td>
<td>1/4th part w/w</td>
</tr>
<tr>
<td>2</td>
<td>Dhav (Anogeissus latifolia)</td>
<td>Bark</td>
<td>1/8th part w/w</td>
</tr>
<tr>
<td>3</td>
<td>Kushta (Saussurea lappa)</td>
<td>Root</td>
<td>1/8th part w/w</td>
</tr>
<tr>
<td>4</td>
<td>Hartala (Arsenic trisulfide)</td>
<td>-</td>
<td>1/8th part w/w</td>
</tr>
<tr>
<td>5</td>
<td>Manshila (Realgar)</td>
<td>-</td>
<td>1/8th part w/w</td>
</tr>
<tr>
<td>6</td>
<td>Haridra (Curcuma longa)</td>
<td>Rhizome</td>
<td>1/4th part w/w</td>
</tr>
<tr>
<td>7</td>
<td>Daruharidra (Berberis aristate)</td>
<td>Stem</td>
<td>1/8th part w/w</td>
</tr>
<tr>
<td>8</td>
<td>Til Tail (Sesamum indicum)</td>
<td>-</td>
<td>1 part w/w</td>
</tr>
<tr>
<td>9</td>
<td>Water</td>
<td>-</td>
<td>4 parts w/w</td>
</tr>
</tbody>
</table>

Table 2. Ingredients and quantity of Aargwadhadi ointment

<table>
<thead>
<tr>
<th>Sr.no</th>
<th>Ingredient</th>
<th>Proportion</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Aargwadhadi Oil</td>
<td>1 part</td>
</tr>
<tr>
<td>2</td>
<td>White wax</td>
<td>1/5th part</td>
</tr>
</tbody>
</table>

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All the procured and authenticated drug of *Aargwadhadi Oil* will be dried and *Kaika* will be prepared.

*Aargwadhadi kaika* (1/4 Part) + *Tila Tail* (1 part) + Water 4 part will be taken in a wide mouth steel vessel.

Above mixture will be heated on *Mandagni* (low flame-gently boiling) with frequent stirring till the *Snehasiddhi Lakshans* appears.

*Tail* will be allowed to get *SwangSheeta* (self-cooled)

After cooling *Tail* will be filtered through doubled muslin cloth and stored in an air tight glass bottle.

Fig. 1. Flow diagram of preparation of *Aargwadhadi Oil*

*Aargwadhadi Oil* will be placed over mild heat until it starts foam appearing

1/5 th part of white wax will be added to the *Aargwadhadi Tail*

Wax will be completely melted in oil, it will be filtered and kept in another vessel

Fig. 2. Preparation of *Aargwadhadi* ointment

2.4 Analytical Study of *Aargwadhadi* Ointment

2.4.1 Organoleptic Characters

*Sparsh* (Touch), *Roop* (Colour, appearance), *Gandha* (Odour)

Physio-chemical Parameters such as pH, Viscosity, Homogeneity, Spreadability Test, Skin Irritation Test, TLC, Microbiologic test, photosensitization will be assessed.

2.5 Experimental Study

2.5.1 Procurement and selection of animals

Required healthy young adult (at least 8-10 weeks old) male/female albino rats will be selected from animal house. Total 30 Wistar strain albino rats weighing between 200-300 gm, having healthy intact skin will be taken randomly for study.

a. Inclusion criteria

- Healthy young Wistar albino mice of commonly used laboratory strains of either sex will be considered
- Rats weighing about 200-250 gm will be included.
- Rats having healthy and intact skin
- Nulliparous.

b. Exclusion criteria

- Pregnant and diseased mice
- Mice which are under trial of other experiment
Table 3. Grouping of animals and dose administration in Acute Dermal Toxicity study

<table>
<thead>
<tr>
<th>Group</th>
<th>Drug</th>
<th>Dose</th>
<th>No. of Animals</th>
<th>Duration</th>
<th>Route</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group I</td>
<td>Control group- No Medicine</td>
<td>-</td>
<td>10 (5 M +5 F)</td>
<td>14 days</td>
<td>-</td>
</tr>
<tr>
<td>Group II</td>
<td>Aargwadhadi Oil</td>
<td>2000 mg/kg</td>
<td>10 (5 M +5 F)</td>
<td>14 days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group III</td>
<td>Aargwadhadi Ointment</td>
<td>2000 mg/kg</td>
<td>10 (5 M +5 F)</td>
<td>14 days</td>
<td>Topical</td>
</tr>
</tbody>
</table>

2.6 Methodology of Sub-acute Dermal Toxicity [20]

Subacute dermal toxicity test will be performed according to the OECD guideline 410 for testing of Aargwadhadi Oil & Aargwadhadi ointment. The test drug will be applied daily to the skin to groups of experimental animals, one dose per group, for a period of 28 days. During the period of application of treatment, the animals will be observed daily to perceive signs of toxicity. Animals, which died during the tests, will be necropsied and at the conclusion of the tests the surviving animals will be sacrificed and necropsied. About 10 % of the body surface area will be cleared for the application of the test drug. Each group contain ten healthy rats (n=10). Based on OECD guidelines 410, at least three doses of Tail and ointment, as well as a vehicle and control groups, will be used in the study. In addition, a satellite group of 10 animals (5 animals per sex) may be treated with the high dose level for 28 days and observed for reversibility, persistence, or delayed occurrence of toxic effects for 14 days post-treatment (Table 4).

2.7 Assessment Criteria

1. General observation
   a) Changes in behavioral Pattern.
   b) General appearance
   - CNS depression
   - CNS stimulation
   - Salivation
   - Diarrhea

   2. Changes in body weight (weekly)
   3. Food and water consumption (weekly)
   4. Fecal consistency (weekly)
   5. Hematology

Qualitative urine examination will be done before exposure to the test compound and the experiment using Uristik A 10 reagent strips at the termination of the trial.

Blood samples will be collected under light ether anesthesia, following hematological values will be analyzed.

- Packed cell Volume (PCV),
- Hemoglobin (Hb)
- Red blood cell (RBC) count,
- Total white blood cell (WBC) count,
- Differential Leucocyte Count (Polymorph and Lymphocyte ratio) and
- Prothrombin Time (PT)
- Plasma Glucose (mg/dl),
- Creatinine (mg/dl),
- SGOT (U/L) and SGPT (U/L)

Blood samples will be withdrawn from the posterior vena cava of each rat and will be collected in EDTA vacuumed blood collection tubes and will gently mixed immediately to mix the blood with EDTA-anticoagulant material inside the tubes for automatic and manual hematological analyses.

6. Biochemical SGOT, SGPT, Alkaline Phosphate, Serum creatinine, Serum urea, Serum Electrolyte, Sodium and Potassium.

2.8 Histopathology Study for Acute Dermal Toxicity Study

As per the need of animals will be sacrificed after 14 days by administering mild anesthesia. Liver, heart, kidney, intestine, stomach, brain will be processed for histopathological studies as per the prescribed procedures.
Table 4. Grouping of animals for Subacute dermal toxicity study (n=10)

<table>
<thead>
<tr>
<th>Group</th>
<th>Drug</th>
<th>Dose</th>
<th>No. of Animals</th>
<th>Duration</th>
<th>Route</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group I</td>
<td>Control group-No Medicine</td>
<td>-</td>
<td>10 (5 M +5 F)</td>
<td>28 Days</td>
<td>-</td>
</tr>
<tr>
<td>Group II</td>
<td>Aargwadhadi Oil</td>
<td>500 mg/kg</td>
<td>10 (5 M +5 F)</td>
<td>28 Days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group III</td>
<td>Aargwadhadi Oil</td>
<td>1000 mg/kg</td>
<td>10 (5 M +5 F)</td>
<td>28 Days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group IV</td>
<td>Aargwadhadi Oil</td>
<td>2000 mg/kg</td>
<td>10 (5 M +5 F)</td>
<td>28 Days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group V</td>
<td>Aargwadhadi Ointment</td>
<td>500 mg/kg</td>
<td>10 (5 M +5 F)</td>
<td>28 Days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group VI</td>
<td>Aargwadhadi Ointment</td>
<td>1000 mg/kg</td>
<td>10 (5 M +5 F)</td>
<td>28 Days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group VII</td>
<td>Aargwadhadi Ointment</td>
<td>2000 mg/kg</td>
<td>10 (5 M +5 F)</td>
<td>28 Days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group VIII</td>
<td>No Medicine Recovery phase</td>
<td>-</td>
<td>Group IV Animal</td>
<td>14 Days</td>
<td>-</td>
</tr>
<tr>
<td>Group IX</td>
<td>No Medicine Recovery phase</td>
<td>-</td>
<td>Group VII Animal</td>
<td>14 days</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 5. Assessment criteria (Erythema) [21]

<table>
<thead>
<tr>
<th>Skin reaction</th>
<th>Grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slight, spotty/diffuse</td>
<td>1</td>
</tr>
<tr>
<td>Moderate uniform redness</td>
<td>2</td>
</tr>
<tr>
<td>Dark red with oedema</td>
<td>3</td>
</tr>
<tr>
<td>Fiery red with oedema or epidermal defect</td>
<td>4</td>
</tr>
</tbody>
</table>

Table 6. Assessment criteria [Scaling]

<table>
<thead>
<tr>
<th>Skin reaction</th>
<th>Grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dryness shiny</td>
<td>1</td>
</tr>
<tr>
<td>Fine scale</td>
<td>2</td>
</tr>
<tr>
<td>Moderate</td>
<td>3</td>
</tr>
<tr>
<td>Severe</td>
<td>4</td>
</tr>
</tbody>
</table>

Table 7. Assessment criteria [Fissures]

<table>
<thead>
<tr>
<th>Skin reaction</th>
<th>Grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fine cracks</td>
<td>1</td>
</tr>
<tr>
<td>Single/multiple broader fissure</td>
<td>2</td>
</tr>
<tr>
<td>Cracks with hemorrhage or exudation</td>
<td>3</td>
</tr>
</tbody>
</table>

Table 8. Assessment criteria [Oedema]

<table>
<thead>
<tr>
<th>Skin reaction</th>
<th>Grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very slight oedema</td>
<td>1</td>
</tr>
<tr>
<td>Slight oedema</td>
<td>2</td>
</tr>
<tr>
<td>Moderate oedema</td>
<td>3</td>
</tr>
<tr>
<td>Severe oedema (raised more than 1mm)</td>
<td>4</td>
</tr>
</tbody>
</table>
2.9 Histopathology Study for Subacute Dermal Toxicity Study

Histopathology will be done by administrating mild anesthesia. Histological examination should be performed on the preserved organs and tissues of the high dose group and the control group. Animals in the satellite group should be examined histologically with particular emphasis on those organs and tissues identified as showing effects in the other treated groups. Liver, kidney, heart, brain, intestine, lungs, stomach will be processed for histopathological studies as per prescribed procedure. Skin, liver, and kidney samples will be collected and fixed in 10% formalin for 48 hrs.

2.10 Rating of SKIN REACTION for the Experiment (Table 5-8)

2.10.1 Methodology of efficacy study of aargwadhadi oil and Aargwadhadi ointment on animal model of vitiligo [22-23]

Experiment will be conducted on C57BL/6 mice of 4 weeks of age. Animals will be divided into five groups consisting of 6 mice in each group.

Dorsal region will be shaved, approximately 24 hr before. Monobenzone 40% cream will be daily applied (50 mL, for 50 days) to the dorsal region (1x1 cm²) the same site near the tail. Cream was massaged until complete absorption using a spatula. In a different dorsal region (near the neck, 1x1 cm²) Aargwadhadi Oil or Aargwadhadi ointment as per grouping (daily, twice a day) will be applied and massaged until complete absorption. All treatments will be carried out till 65 days. 6 mm circles of ear tissue will be collected. All tail and back skin will be removed.

Some samples will be stored at 80°C for further tests. Same dorsal skin samples will be collected and placed in 10% neutral buffered formalin to histological analysis. Samples will be sectioned and stained with Fontana Masson and then photograph in increments of 200x will be taken, analysis will be performed by counting cell per field. Ten fields from three histological sections of each group will be analyzed (Table 9).

2.11 De-pigmentation Evaluation

The extent of de-pigmentation will be analyzed by objective observation by two blinded observers.

2.12 Type of Study

Pharmaceutical, Analytical, and Experimental Interventional Animal Study.

2.13 Study Duration

3 months

2.14 Study Centers

1. Dattatraya Ayurved Rasashala, Mahatma Gandhi Ayurved College Hospital and Research Center, Salod (H) Wardha.
2. Animal house, Datta Meghe Pharmacy College, DMIMS(DU), Salod (H) Wardha
3. Analysis of the formulation will be done at Central Research Lab, Jawaharlal Nehru Medical College, Sawangi (Wardha).
4. According to need, study will be carried out at certified or standard institute / organization/ lab recognized or recommended by DMIMS (DU).

<table>
<thead>
<tr>
<th>Group</th>
<th>Drug</th>
<th>Dose</th>
<th>No. of Animals</th>
<th>Duration</th>
<th>Route</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group I</td>
<td>Untreated group no medicine for monobenzone induced animal sample</td>
<td>-</td>
<td>6[3M+3F]</td>
<td>50days</td>
<td>-</td>
</tr>
<tr>
<td>Group II</td>
<td>Oil/Vehicle control group</td>
<td>2000mg/kg Til tail</td>
<td>6 (3 M +3F)</td>
<td>50 days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group III</td>
<td>Ointment Vehicle control group</td>
<td>2000 mg/kg White paraffine cream</td>
<td>6(3 M +3 F)</td>
<td>50 days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group IV</td>
<td>Experimental group</td>
<td>2000 mg/kg Aargwadhadi oil</td>
<td>6[3M+3F]</td>
<td>50 Days</td>
<td>Topical</td>
</tr>
<tr>
<td>Group V</td>
<td>Experimental group</td>
<td>2000 mg/kg Aargwadhadi ointment</td>
<td>6[3m+3F]</td>
<td>14 Days</td>
<td>Topical</td>
</tr>
</tbody>
</table>
3. OBSERVATION AND RESULTS

- Observations will be noted and presented in the form of tables, chart, photographs etc.
- The data will be analyzed with application of suitable inferential statistics

3.1 Method of Data Analysis Statistical Analysis

The mean and standard deviation of the treated groups will be done by applying unpaired t-test and ANOVA analysis. The data obtained will be statistically analyzed by using Statistical Package for statistical Science (SPSS) software version 23. The data generated would be mentioned as Mean ± SEM difference among the groups would be assessed by employing student ‘t’ test and ANOVA analysis.

4. DISCUSSION

The basic Ayurvedic principles of ease of availability (natural abundance), cost effectiveness, and ease of topical application keeping in consideration the demands of current lifestyle trends and client preferences will be followed to provide a holistic remedial solution to varied issues of vitiligo and skin diseases. This formulation has been devised in accordance with the overarching guiding principles of Ayurveda that encourage individual preference and ease of preparation and application. Considering current lifestyle and trending preference amongst users, a standardized formulation in both, oil and ointment form with safety is the most important consideration before administration of any herbal product in human being. In However, herbal medicines are not completely safe from adverse effects if not used judiciously [24-25]. So acute and subacute dermal toxicity will be beneficial for providing safe product for skin disorders especially for vitiligo.

The ointment form is easier to carry and apply, and also has a better absorption [26]. The efficacy of the drug will be tested by comparing the effect of conventional Aargwadhadi oil and the Aargwadhadi ointment formulation for its efficacy in vitiligo. The dosage and frequency will be suggested at the outset and following experimental, the researcher expects to establish the efficacy, acceptability and outcome of the Aargwadhadi oil and Aargwadhadi ointment. Studies on efficacy of different Ayurvedic preparations were reported [27] but the results are not encouraging. Hence present formulation may show a ray of hope in the treatment of vitiligo.

5. CONCLUSION

The present available treatment for Vitiligo does not provide complete remission of symptoms for long duration. Thus the newly developed herbal formulation will provide significant result in controlling Vitiligo and modified dosage i.e. ointment will overcome the limitation of oil and provide a cost effective cosmetic product in controlling Vitiligo.

6. STRENGTH OF STUDY

Study will create standard operating procedure about preparation and analytical studies of Aargwadhadi ointment.

Research formulation Aargwadhadi Oil can be converted in to new dosage form such as Cream which is convenient for external application.

DISCLAIMER

The products used for this research are commonly and predominantly use products in our area of research and country. There is absolutely no conflict of interest between the authors and producers of the products because we do not intend to use these products as an avenue for any litigation but for the advancement of knowledge. Also, the research was not funded by the producing company rather it was funded by personal efforts of the authors.

NOTE

The study highlights the efficacy of "Ayurveda" which is an ancient tradition, used in some parts of India. This ancient concept should be carefully evaluated in the light of modern medical science and can be utilized partially if found suitable.

CONSENT AND ETHICAL APPROVAL

This will be taken from institutional ethical committee of Datta Meghe Institute of Medical Sciences, Sawangi, (DMIMS) Wardha and from Animal ethics committee. Study will be followed as per instructions of IAEC of DMIMS. As per international standard or university standard, respondents’ written consent will be collected and preserved by the author(s).
COMPETING INTERESTS

Authors have declared that no competing interests exist.

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