Pharmacognostical Standardisation and HPTLC Quantification of Gallic acid in Homonoia riparia Lour

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INTRODUCTION

Homonoia riparia Lour. (Family: Euphorbiaceae) commonly known as Kshudra Pashanbhed in the Indian system of medicine grows widely across Asian countries and is planted along rivers and streams to stabilize and protect the banks.1 Historically, the plant has been used in Indian and Chinese systems of medicine for the treatment of various ailments. In Cambodia, stem and leaves are used as purgative where as wood in the heart wood and bark of Homonoia riparia Lour. is widely used in Indian and Chinese system of medicine. Microbiological action was given in malaria and scabies. Leaves are used in Malaysia and Thailand for skin infections. Traditionally, root of the plant is used in India as a laxative, emetic, diuretic, to treat urinary calculi, syphilis, gonorrhoea, malaria, antiseptic, inflammation, ulcers, urinary discharges, urinary infection, hepatitis and fungal infections.2-4 Six new cycloartane type triterpenes and quercetin glycoside have been isolated from the leaves of Homonoia riparia. β-sitosteryl, stigmasterol, taraxerone, taxe rol, lupenol, gallic acid, β-acetyl aleuritolic acid have been isolated from the heart wood and bark of Homonoia riparia.5,6 In addition, riparsaponine (cycloartane-type triterpenoid) was also reported from the stem of Homonoia riparia.7 Taking into consideration the medicinal potential of this plant, aim of the present study was to establish pharmacognostical and phytochemical standardisation parameters for the plant Homonoia riparia and to identify potential marker phytoconstituents in Homonoia riparia that could serve as reference standards.

MATERIALS AND METHODS

Chemicals

Analytical grade chemicals such as safranin, formalin, chloral hydrate, acetic acid, ethyl alcohol, glycerine, iodine, potassium iodide, phloroglucinol, and hydrochloric acid were used for the present study. Gallic acid was purchased from Sigma Chemicals Co, USA. HPTLC plates were obtained from E. Merck, Germany.

Plant material

Homonoia riparia was procured from the shores of Sitanadi, Karnataka in the month of February and authenticated by Dr. Gopala Krishna Bhat, Professor (Rtd.), Poorna Prajna College; Udupi and voucher specimen was procured from the shores of Sitanadi, Karnataka

ABBREVIATIONS AND ACRONYMS


PICTORIAL ABSTRACT

PICTORIAL ABSTRACT

has been submitted in the herbarium of Manipal college of Pharmaceutical Sciences, Manipal. The plant material was coarsely powdered and stored in a well closed container for phytochemical analysis and determination of pharmacopoeia standards.

**Preparation of methanol extract**
Powdered plant material was extracted with methanol using Soxhlet extraction unit and concentrated under rotary evaporator to yield a brownish green coloured mass and was stored in refrigerator for phytochemical and High Performance Thin Layer Chromatographic analysis.

**Macroscopic characteristics**
The macroscopical features such as shape, texture, colour, odour and taste of the whole plant of *Homonoia riparia* were determined.

**Microscopic evaluation**
Thin transverse section of root, stem and leaf were taken using a razor blade and stained with safranin as per standard procedures and anatomical characters were distinguished under Zeiss AXIO trinocular microscope. Photographs of sections were taken using Zeiss Axio Cam attached with the microscope under bright field light.

**Powder microscopy**
The whole plant material was powdered and passed through sieve number 60 and treated with phloroglucinol and concentrated hydrochloric acid and various cell characters were observed under microscope.

**Physicochemical evaluation**
The whole plant of *Homonoia riparia* was subjected to various physicochemical evaluation methods such as total ash, acid insoluble ash, water soluble ash, extractive value (water and ethanol), moisture content, foaming index and swelling index as per WHO guidelines.

**Fluorescence analysis**
Fluorescence analysis of whole plant of *Homonoia riparia* was carried out under visible and UV light (266 nm and 366 nm) after treatment with various chemical reagents.

**Phytochemical analysis**
Phytochemical analysis of methanol extract of *Homonoia riparia* was performed using different chemical tests in order to check the presence of secondary metabolites such as sterols, steroids, fatty acid, alkaloids, glycosides, flavonoid, and tannins in the plant by standard methods.

**Quantification of gallic acid by HPTLC analysis**
CAMAG High Performance Thin Layer Chromatography (HPTLC) instrument was used for the quantification of gallic acid in the methanol extract of *Homonoia riparia*. The standard solution was prepared at a concentration of 100 µg/mL and test solution at 5 mg/mL. Ten microlitres of the sample was applied as 8 mm bands on pre-coated silica gel plate using Camag Linomat 5 applicator in duplicate. The HPTLC plates were developed in solvent system, toluene: ethyl acetate: formic acid: methanol (6:6:1:80.25 v/v). After development, plate was dried and scanned at 280 nm using CAMAG TLC scanner 3. The peak areas were recorded and percentage content of gallic acid was determined.

**Statistical analysis**
The experiments were carried out in triplicate and the results are reported as mean ± standard error of mean (SEM).

## RESULTS

**Macroscopic characteristics**
The roots arise from a common root stock and are hard, cylindrical, 5-30 cm long, brown in colour with attached rootlets and fibrous fracture. It has no specific odour and taste. Leaves are dorsiventral, simple, alternate and linear to lanceolate shaped, 14-22 cm long, and 1.6 to 2 cm wide. The upper surface is greenish in colour and shining whereas lower part of leaf is light green in colour and hairy. The leaves show pinnate venation, acute apex, cuneate base and entire margin. Stem is hard and nodes and internodes are present. The outer surface is brownish and inner surface light yellow in colour with no characteristic odour and taste. The fracture is fibrous (Figure 1).

**Microscopic characteristics**

### Root
Transverse section of the root shows cork, cortex and stellar regions (Figure 2a). Cork region consist of 3-5 layers of tangentially elongated cells with deposition of brown coloured matter. Cork cambium (phellogen) is one layer thick and indistinct. Cortex consists of several layers of parenchymatous cells which are polyhedral. Group of pericyclic fibres are also present in this region (Figure 2b). Secondary xylem is chiefly made of rounded vessels, thick walled parenchyma and highly lignified fibres. Concentric starch grains are distributed in the xylem region. Radially arranged non-lignified medulary rays are visible in the phloem region and that in xylem region are lignified. Thick walled parenchymatous cells are present in the pith region with high deposition of starch grains (Figure 2c).

### Transverse section of leaf
Transverse section of leaf consists of lamina and midrib regions. The lamina shows upper and lower epidermis; rectangular epidermal cells, and anomocytic stomata. Mesophyll comprises of palisade and spongy parenchyma cells filled with chloroplast. Rosette crystals are present. Palisade cells are single layered and they are cylindrical and spongy parenchyma has four layers of spherical or loosely arranged cells. Midrib exhibit crescent shaped bicollateral vascular bundle ensheathed by sclerenchymatous bundle sheath. The parenchymatous ground tissue lies...
Figure 2: (A): Microscopy of root of *Homonoia riparia* shows Ck-cork; Ct-Cortex; XY-xylem. (B): Transverse section of the xylem region of root showing the presence of SG-Starch grains, Ve-Xylem vessels; XR-xylem rays; XF-Xylem fibres. (C): Enlarged portion of cork and cortex region of root showing CC-Cork cambium; MR-Medullary rays; PF–Phloem fibres; Ct-Cortex; RC-Rosette crystal.

Figure 3: (A) Microscopy of leaf of *Homonoia riparia* showing. F–fibre; LE–lower epidermis; Me–mesophyll; Pal–palisade; Ph–phloem; RC-rosette crystal; SP–spongy parenchyma; UE–upper epidermis; XY–xylem. (B) T.S of midrib showing. F–fibre; LE–Lower epidermis; Ph–Phloem; RC-Rosette crystal; VB–Vascular bundles; XY–Xylem. (C) Transverse section of stem showing. Ck–cork; Ct–cortex; F–fibre; Ph–phloem; XY–xylem fibres; Pi–pith. (D) Microscopy of cork and cortex region of stem showing Ck–Cork; CC-Cork cambium; Scl–Sclereids; RC-Rosette crystal; F–fibre.
next to bundle sheath. Xylem and phloem are the components of the vascular bundle. Starch grains are absent. (Figures 3a and 3b).

**Stem**

Transverse section of mature stem shows phellem consisting of 3-5 layers of tangentially elongated cork cells with highly suberised walls. The phellogen (cork cambium) is single layered consisting of radially elongated thin walled cells. Phellogen is followed by phelloderm and cortex, indistinguishable from each other; consisting of 10-12 layered loosely thin walled parenchymatous cell. Stone cells, a reduced form of sclerenchyma cells appears either solitary or in groups of two to three cells, pitted and highly thickened. Cortex is followed by secondary phloem consisting of phloem parenchyma, phloem fibres with narrow lumen, and uniseriate medullary rays. Secondary cortex also contains rosette calcium oxalate crystals. Pith region consists of parenchymatous cells and group of pericyclic fibres. Simple, granular starch grains are present in the phloem, xylem, ray and pith region (Figures 3c and 3d).

**Powder microscopy**

Powdered drug of *Homonoia riparia* was yellowish brown, odourless and coarse in texture. Microscopically, the powder showed lignified fibres, cork cells, spirally annulated xylem vessels and anomocytic stomata (Figure 4).

![Figure 4: Powdered characteristics of the whole plant of Homonoia riparia showing A) St-Anomocytic stomata; B) XV-Spiral xylem vessels; C) Lignified fibres D) Ck- Cork cells.](image)

**Physicochemical evaluation**

Ash value determination is useful for identifying low quality product, earthy matter or excess of sandy matter or impurities present with drug. The percentage of total ash, acid insoluble ash and water soluble ash were calculated. The results are shown in Table 1. Extractive values are useful to indicate the nature of chemical constituents and also for the identification of adulteration. The extractive values (ethanol and water) are presented in Table 2. The moisture content, swelling index and foaming index were calculated and shown in Table 3. The fluorescence analysis of sample drug is tabulated in Table 4.

**Table 1: Ash values of Homonoia riparia**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Value % w/w</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total ash</td>
<td>5.02 ± 0.59</td>
</tr>
<tr>
<td>Water soluble ash</td>
<td>3.38 ± 0.05</td>
</tr>
<tr>
<td>Alcohol insoluble ash</td>
<td>1.175 ± 0.74</td>
</tr>
</tbody>
</table>

**Table 2: Extractive values of Homonoia riparia**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Value % w/w</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol soluble extractive value</td>
<td>6.93 ± 0.01</td>
</tr>
<tr>
<td>Water soluble extractive value</td>
<td>11.4 ± 0.02</td>
</tr>
</tbody>
</table>

**Table 3: Moisture content, foaming index and swelling index of Homonoia riparia**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture Content</td>
<td>6 ± 0.61 (w/w)</td>
</tr>
<tr>
<td>Foaming Index</td>
<td>111.11 ± 0.3</td>
</tr>
<tr>
<td>Swelling Index</td>
<td>4 ± 0.1</td>
</tr>
</tbody>
</table>
Phytochemical analysis

Methanol extract of *Homonoia riparia* indicated the presence of primary and secondary metabolites such as carbohydrates, sterols, flavonoids, phenolic compounds, tannins, saponins etc (Table 5).

**Table 5: Preliminary phytochemical screening of *Homonoia riparia***

<table>
<thead>
<tr>
<th>Phytoconstituents</th>
<th>Methanol extract</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkaloids</td>
<td>-</td>
</tr>
<tr>
<td>Carbohydrates</td>
<td>+</td>
</tr>
<tr>
<td>Sterols and triterpenoids</td>
<td>+</td>
</tr>
<tr>
<td>Saponins</td>
<td>+</td>
</tr>
<tr>
<td>Fixed oils and fats</td>
<td>-</td>
</tr>
<tr>
<td>Phenolic compounds and tannins</td>
<td>+</td>
</tr>
<tr>
<td>Proteins and amino acids</td>
<td>-</td>
</tr>
<tr>
<td>Flavonoids</td>
<td>+</td>
</tr>
</tbody>
</table>

Quantification of gallic acid by HPTLC

Presence of gallic acid in the methanol extract of *Homonoia riparia* was confirmed by HPTLC analysis and the percentage of gallic acid (R_f 0.56) was found to be 0.76% w/w (Figure 5).

**DISCUSSION**

Quality control standards of herbal medicine used in indigenous systems of medicine have become highly applicable due to the reawakening of interest in these drugs all over the world. In the present study, the morphological, microscopical, physicochemical and phytochemical characteristics of the whole plant of *Homonoia riparia* have been reported. Morphological and microscopical parameters were established to authenticate the genuine drug. These features will be useful for easy and quick identification of the right variety and adulterants. The physical evaluation of the crude drugs plays an important criterion in detecting adulteration or improper handling of drugs. Physicochemical reports showed that total ash value was 5.02% w/w, acid insoluble ash was 1.18% w/w, water insoluble ash was 3.38% w/w, foreign organic matter 0.7% w/w, and alcohol soluble extractive value was 6.93% w/w and water soluble extractive value was 11.4% w/w. The moisture content of drug was found to be considerably high (Table 3). Lower moisture content indicates better stability against decomposition of the drug and it should be reduced to avoid decaying of the crud drug by chemical change or microbial attacks. The results of the phytochemical screening of whole plant of *Homonoia riparia* indicated the presence of terpenoids, flavonoids, phytosterols, phenolic compounds, tannins, saponins which might show therapeutic potential in further biological investigations. Gallic acid and its structurally related compounds are biologically active phytoconstitu...
ents widely seen in fruits and plants and are found to possess number of pharmacological activities such as anticancer, anti-inflammatory, cardioprotective, hepatoprotective, and renoprotective activities. Marker compound based standardization has been demonstrated to be a valid method for the identification of botanical medicine and formulations. The percentage of marker compounds serve as an indicator of the quality of plant materials. In the present study, significant quantity of gallic acid was detected and quantified in the methanol extract of Homonoia riparia that could act as a marker for standardization. Till date, there are no reports of the pharmacognostical standardization of Homonoia riparia, therefore, our study will provide significant information for identification and authentication of plant material and serve as a quality control for future reference.

CONCLUSION

The present study was carried out to establish the detailed pharmacognostical and physico-phytochemical parameters of the plant Homonoia riparia that will aid in the correct authentication and identification of the plant material.

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CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

REFERENCES


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