PHARMACOCGNOSTIC AND PHYTOCHEMICAL EVALUATION OF CARYOTA URENS LEAF

Mariyan R. Patel *, Hiteksha S. Panchal 1, Ajay K. Saluja 2
1Indubhai Patel Pharmacy College & Research Center, Dharmaj, Virsad, Gujarat, India
2A R College of Pharmacy, V.V. Nagar, Anand, Vidyanagar, Gujarat, India
*Corresponding Author Email: mary.patel20@gmail.com

ABSTRACT

Caryota urens (Palmae) is native to India. This tree species has been of interest to researchers because it is a medicinal plant employed in the Indian indigenous system of medicine. Pharmacognostic standardization, physico-chemical evaluation of the leaves of Caryota urens was carried out to determine its macro and micro-scopical characters and also some insoluble ash and sulphated ash values, alcohol- and water-soluble extractive values were determined for phytochemical evaluations. Preliminary phytochemical screening was also done to detect different phytoconstituents. Microscopically, Leaf showed Lamina, midrib regions, stomata. Powder microscopy showed mesophyll region, pitted xylem vessels and paracytic stomata. TLC of petroleum ether and ethanol extract showed three spots using Hexane: Ethyl acetate (12:4) and three spot using Chloroform: Ethyl acetate (5:4). Phytochemically leaves exhibited phytosterols, Flavonoids, Tannin carbohydrates and phenolic compounds.

Keywords: Caryota urens, macroscopy, microscopy, phytoconstituents, Rf (retention factors)

INTRODUCTION

Caryota urens is popularly known as Shankarjata in Ayurveda is distributed more or less throughout Sub Himalaya tract from Nepal eastwards, Very common in Ghats and sub montane forests of Mysore, in Malabar, Coorg, Konkan, Cochin and Travancore. Also in tropical Asia, Malaya, Singapore, Australia and Ceylon. 1,2 Leaf is reported to cure hemicrania 3 and Root of the plant is reported for treatment of abortion,dysentery and tooth cavity to prevent decay 4. Literature revealed that pharmacognostic studies have not been reported for the leaf of this plant. Therefore the main aim of the present work is to study the macro, microscopic and some other pharmacognostic characters and physico-chemical standards of leaf of Caryota urens which could be used to explore this plant.

MATERIALS AND METHODS

Collection of plant material: The plant Caryota urens was authenticated by Dr. Bhanu Kakrani, Lecturer, Dept. of Botany, Tolani College of Arts & Science, Adipur (Kutch) .Voucher herbarium specimen [HSP/TO-6/37] is preserved along with crude drug sample at the herbarium of A. R. College of Pharmacy, Vallabh Vidyanagar.

Pharmacognostic evaluation

Macroscopy: The macroscopic characters such as color, odour, taste, nature, texture were studied for morphological investigation. The shape, apex, base, margin, taste and odour were determined in case of leaves 5.

Microscopy 6

Preparation of transverse section of leaf and stem: The T.S. of leaf and stem were cleared of coloring matter by heating with chloral hydrate. After clearing, the set of slides were mounted in glycerin. Another set of sections was stained with phloroglucinol and concentrated hydrochloric acid (1:1) mixture for differentiating lignified tissues. A third set of sections was treated with dilute iodine solution. All the sections were then observed under 10X and 45X.

Surface preparation of the leaf: Small portion of leaf (2 mm square) was placed in chloral hydrate solution in test tube; epidermis was exposed by scraping of the tissues with sharp edge of razor on the glass slide. Water was added slowly and continuously scraping was done till transparent. The portion was mounted in a mixture of equal portion of glycerin.

Powder preparation of leaves and stem: The aerial parts of Caryota urens were dried under shade. The plant parts were powdered by grinding and passed through the sieve number 60. Finally, from this coarse powder, microscopic examination was done. Slides were prepared in same manner as mentioned in above method.

Quantitative Microscopy 7: Quantitative leaf microscopy to determine palisade ratio, stomatal number, stomatal index, palisade ratio, vein islet number and vein termination number were carried out on epidermal strips. Other quantitative microscopic parameters are determination of fibre size in powder of aerial parts.

Proximate Analysis 8,9,10: The ash values, extractive values, moisture content, crude fibre content, swelling index, elemental analysis, foaming index were performed according to the official methods prescribed in Indian Pharmacopoeia and the WHO guidelines on quality control methods for medicinal plant material

Phytochemical Analysis: The dried powdered plant material was successively extracted with the solvents of increasing polarity in a Soxhlet apparatus utilizing petroleum ether (60 - 80), Toluene, acetone, chloroform, methanol and water. The liquid extracts obtained with different solvents were collected and the consistency, color, appearance of the dried extracts and their percentage yield were noted. The extracts obtained from powder by successive solvent
Physico-chemical evaluations

Physicochemical parameters of *Caryota urens* leaf powder were determined and reported as total ash, water-soluble ash, acid-insoluble ash, and sulphated ash values. Alcohol and water-soluble extractive values were determined to find out the amount of water and alcohol soluble components\textsuperscript{14}. The moisture content was also determined. As a part of quantitative microscopy stomatal number, stomatal index and fibre size were determined by using fresh leaves of the plant.

RESULT AND DISCUSSION

Macroscopy

Leaf is 24-28 cm × 6-8 cm, triangular, Dark to light Green color, Premorse apex, Irregularly serrate; equal sided at the base; smooth surface both side, venation parallel ;[Figure 1] Taste: sweet ; Odour: without characteristic.

Microscopy

A Transverse Section of leaf shows lamina and midrib portion. Leaf is isobilateral type. It is not clearly differentiated into palisade and spongy parenchyma. Upper and lower epidermis (Ep) having no trichomes. Upper and lower epidermis both are double layered. Midrib portion is well developed with lignified parenchyma in which some parenchymas are pitted. In the middle region presence of vascular bundles in distributed position and some tracheid are also there.

Surface preparation of leaf

In the surface view of the leaf, the lower epidermal cells having more no of stomata than in upper epidermis. It shows paracytic type stomata.
Powder microscopy of dried powder of leaf

The powder of *C. urens* was greenish color, without characteristic odour and with sweet taste. When powder was mounted with chloral hydrate, phloroglucinol and HCl the following elements were observed: Mesophyll region which contain palisade with epidermis, paracytic stomata [Figure 4].

![Paracytic stomata](image1)

![Peicyclic fibre](image2)

![Pi Xy: Pitted Xylem vessel](image3)

Figure 4: Powder Study of leaf of *Caryota urens*

Physico-chemical evaluation

<table>
<thead>
<tr>
<th>Sr. no.</th>
<th>Parameter</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Stomatal number</td>
<td>Lower epidermis: 275</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Upper epidermis : 50</td>
</tr>
<tr>
<td>2.</td>
<td>Stomatal index</td>
<td>Lower epidermis: 14.66</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Upper epidermis: 4</td>
</tr>
<tr>
<td>3.</td>
<td>Fibre size</td>
<td>Length: 145.19 μm - 359.54 μm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Width: 3 μm – 7 μm</td>
</tr>
<tr>
<td>4.</td>
<td>Total ash</td>
<td>19.37 %w/w</td>
</tr>
<tr>
<td>5.</td>
<td>Water-soluble ash</td>
<td>4.75% w/w</td>
</tr>
<tr>
<td>6.</td>
<td>Acid-insoluble ash</td>
<td>15% w/w</td>
</tr>
<tr>
<td>7.</td>
<td>Water-soluble extractive value</td>
<td>17.04% w/w</td>
</tr>
<tr>
<td>8.</td>
<td>Alcohol-soluble extractive value</td>
<td>15.94% w/w</td>
</tr>
<tr>
<td>9.</td>
<td>Moisture content</td>
<td>3.29% w/w</td>
</tr>
</tbody>
</table>

Table 1: Physico-chemical evaluations of Leaf of *Caryota urens*

PHOTOCHEMICAL ANALYSIS

Preliminary profiles of Successive solvent extracts

*C. urens* leaf showed presence of moisture content- 3.29% w/w; total ash, acid insoluble ash and water-soluble ash determined were 19.37, 15 and 4.75% w/w, respectively. Water-soluble extractive value was 17.04; alcohol-soluble extractive value was 15.94; the color consistency and % yield of successive extractive values of powder were petroleum ether (60-80°C) (yellowish brown, sticky mass, 4.7), toluene (Brownish black, sticky mass, 2.53), chloroform (Brownish black, sticky mass, 1.49), ethanol (Brownish black, sticky mass, 13.23), water (Dark brown, solid mass, 4.82) [Table 2].

<table>
<thead>
<tr>
<th>Sr.No.</th>
<th>Solvent</th>
<th>Color and consistency after drying</th>
<th>Average value (%w/w)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Petroleum ether (60-80°C)</td>
<td>Greenish sticky mass</td>
<td>9.044 %</td>
</tr>
<tr>
<td>2.</td>
<td>Toluene</td>
<td>Greenish solid mass</td>
<td>4.125 %</td>
</tr>
<tr>
<td>3.</td>
<td>Chloroform</td>
<td>Greenish solid mass</td>
<td>2.91 %</td>
</tr>
<tr>
<td>4.</td>
<td>Methanol</td>
<td>Yellowish green sticky mass</td>
<td>11.5 %</td>
</tr>
<tr>
<td>5.</td>
<td>Water</td>
<td>Reddish brown, solid mass</td>
<td>9.775 %</td>
</tr>
</tbody>
</table>

Table 2: Preliminary Profile of Successive solvent extracts of Leaves of *Caryota urens*

Preliminary phytochemical screening

All the above extracts were tested with various reagents and the results for the same are reported in table no.3. The various extracts showed the presence of phytosterols, carbohydrates, phenolic compounds, flavonoids, tannins, proteins, amino acids and mucilage.
CONCLUSION

As there is no pharmacognostical anatomical work on records for this traditionally much valued shrub, present work is taken up in the view to lay down the macroscopic and microscopic standards, which could be used in deciding the genuineness of the herb, irrespective of their collection from different sources. The colored photographs of the Leaf of the above mentioned plant might facilitate the researcher for identification. The results of the phytochemical screening, histochemical tests can be considered as distinguishing parameters to identify and decide the authenticity of C. urens and thus can be used as standards for reference purpose also. The outcome of the quantitative parameters described on the above-mentioned plant parts (Leaf) might be useful in determining the authenticity of the herbs. HPTLC profile helps in standardization and also for undertaking work on isolating and identifying the bioactive compounds.

ACKNOWLEDGEMENT

The authors are thankful to the Principal, A. R. College of Pharmacy for evincing interest in this work. They are also thankful to Dr. Bhanu Kakrani, Lecturer, Dept. of Botany, Tolani College of Arts & Science, Adipur (Kutch) for help in providing authentic plant material.

REFERENCES


Cite this article as:

Source of support: Nil, Conflict of interest: None Declared

Disclaimer: IRJP is solely owned by Moksha Publishing House - A non-profit publishing house, dedicated to publish quality research, while every effort has been taken to verify the accuracy of the content published in our Journal. IRJP cannot accept any responsibility or liability for the site content and articles published. The views expressed in articles by our contributing authors are not necessarily those of IRJP editor or editorial board members.