Preparation and physico-chemical standardization of Viburnum punctatum Asava

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ABSTRACT

The leaves, stem and the roots of V. punctatum were collected from Nilgiri hills, Tamil Nadu, India. A primary organic analysis conducted on the species revealed that the presence of bioactive molecules such as tannins, saponins, phenolic compounds (flavonoids) and other phenolic glycosides as their principal phyto-constituents. The crude drug (Patha) was formulated into an asava using conventional anaerobic fermentation process for about 60 days. The formulation was standardized by some 13 methods of physico-chemical analysis to obtain a consistent and reproducible parameters. This study can help in re-evaluating the formulation as well as in conducting further investigation on it.

Keywords: Viburnum, Asava, Patha, viscosity, sugar content.

INTRODUCTION

Viburnum Linn. Species contain sterols, sesqui and triterpenoids, phenolic compounds and their glycosides as their common chemical constituents[1-5]. A few species among 17 in India, namely: Viburnum punctatum Buch.-Ham.ex D.Don, Viburnum coriaceum Blume and Viburnum erubescens Wall.ex DC; have been reported in literature to possess uterine sedative, anti-asthmatic, astringent, anti-inflammatory and anti-microbial activities[6,7]. A verbal enquiry to the local community and plant vendors of Ooty and Coimbatore, Tamilnadu, also supported that the above listed pharmacological activities were traditional and were promising with roots, stem barks and leaves of these species[8].

Among the above listed chemical constituents, phenolic compounds, terpenoids and their glycosides may be the cause for biological responses. In addition to this, a qualitative chemical
screening and spectrophotometric analysis of extracts were performed to reveal that the stem part of these three species contains an appreciable amount and a wide range of phenolic compounds[9,10].

Radical scavenging activities of phenolic compounds play a key role in ameliorating healing and even preventing several ailments in living being. It is a well known fact that the plants synthesis phenolic compounds for diverse purposes, which may be of protective, functional or as metabolic end products in nature[11]. But, human exploit them as valuable medicines/ phyto-pharmaceuticals by focusing on their anti-oxidant potential with or without modification.

A quest for a search of herbal phenolic compounds is still a renewed interest in the science of natural products as a source of valuable medicines. The herbal phenolic molecules such as flavonoids, anthocyanins, bioflavones and other phenolic glycosides have, already, been explored and known for their applications against several human ailments-cardiovascular disorders, chronic inflammation and GIT related troubles[12-14]. It is essential to formulate such plants which will be useful in future as a reference.

**MATERIALS AND METHODS**

**Collection of Specimens**
The studies were undertaken on some three parts of the species of the genus *Viburnum* namely: *Viburnum punctatum* Buch.-Ham.ex D.Don. The choices of plant parts were the leaves, stems and the roots of these species. The plant specimens for the study were collected from Nilgiri Hills, Tamil Nadu, India, and authenticated by Dr V Chelladurai, former Professor of Botany, Medicinal Plant Survey for Siddha, Government of India, as *Viburnum punctatum* Buch.-Ham.ex D.Don., (V181) and deposited in the department of pharmacognosy at Nandini Nagar Mahavidyalaya College of Pharmacy, Uttar Pradesh. A care was taken to select healthy plants and the plant parts for the study were collected fresh and dried for a couple of weeks to be involved for further studies.

**Preparation of Viburnum asava by anaerobic fermentation method (An Ayurvedic formulation)**
Approximately 1¼ seers (45 g) of the (patha) were coarsely powdered and added with 32 seers (1024 ml of water) and boiled for about 3 – 5 h. The whole mixture was cooled at room temperature[15]. The mixture was taken in wooden vats of 2 litre capacity, to which dissolved were 12½ seers (400 g) of jaggery and boiled for half an hour.

Dravyas and Dhataki pushpa (*Woodfordia fruticosa*) were then added to the mixture kept in the wooden vats. The vessel was closed with a clean lid followed by wrapping around the lid with seven consecutive layers of clay smeared cloth. The vessel was buried in cellar (basement) for about a couple of months towards the completion of fermentation process (sandhana)[16,17].

After the stipulated meriod (60 days), the vessel was withdrawn to examine the preparation which showed a brownish black fluid with a frothing andaromatic odour and alcoholic taste. The final fluid decanted and filtered through a cotton cloth to obtain a clean transparent asava. Then the asava was bottled and labelled and subjected to some modern methods of standardization.
Standardization of asava [18-21]

Determination of total solids
A shallow, flat bottomed flanged dish, about 75 mm in diameter and about 25 mm deep, made of nickel was used for this analysis. Accurately 5 ml of asava was pipetted out and placed in the dish and evaporated at as low temperature as possible on a water bath until the solvent was removed and the residue is apparently dry. Then the dish was placed in an oven and dried to constant weight at 105°C. After the dish was provided with well-fitting cover, it was cooled in a desiccators.

Determination of boiling range (Distilling range)
A distillation unit fit with a thermometer was employed to determine the boiling range of the asava. The apparatus consisted of a distilling flask of 200 ml capacity; a condenser of 60 cm long; a receiver of 100 ml capacity which was graduated with 1 ml division; and a thermometer showing 0°C - 240°C.

The thermometer was positioned in the centre of the neck and the entire assembly was shield after dropping about 100 ml of asava to the distilling flask. With the aid of metallic stand and clamps, the entire assembly was placed on an electric heater having a thermostat, so that adjustment in temperature could be done conveniently. Distillation was switched on and the recorded was the temperature of first drop of the distillate. Then the temperature was increased in such a way the receiver could collect 4 – 5 ml per min. The process was continued until 25% (25 ml) of the distillate reached the receiver and the temperature of the last drop of the distillate to the receiver was also noted.

Necessary correction was employed observing the temperature readings from any variation in the parametric pressure from the normal (101.3 kPa) using following expression.

\[ t_1 = t_2 + K (a \text{ – } b) \]

\( t_1 \) – corrected temperature; \( t_2 \) – the observed temperature; \( a = 101.3; b \) – the barometric pressure of the time of the determination; \( K \) – the correction factor.

Determination of congealing range or temperature
The congealing temperature is that point at which there exists a mixture of the liquid phase of a substance and a larger proportion of the solid phase. This experimentation required 1 litre beaker in which two test tubes were placed in such a way one was inserted in to another test tube. The inner test tube contained 15 ml of asava and stopperd with a cork attached with a stirrer and a thermometer with 0.2° C graduation.

The beaker was filled with water and the test tubes were clamped in such a way they were immersed in water and distance of 18 mm be maintained between the bottoms of the beaker and test tube. The temperature at which a substance solidifies upon cooling is a useful index of purity.

Preparation of reference substance
Since asava is a liquid, the process of determination of congealing point was carried out in the same way of raising temperature, while stirring, about the room temperature using the apparatus for congealing point determination and noted down as a reference value.
Preparation of test substance of asava
The temperature of the bath was maintained near 15°C using addition of ice cubes and placed on a heating mantle which was kept turned off. Then the sample was stirred constantly to a rate of 20 cycles per min with simultaneous observation of rise in temperature with the thermometer. The congealing point was still hidden up to the room temperature. Hence, a slow rise of temperature was aided to the bath using the heating mantle until the congealing point appeared which was comparable to that of the standard. The process was repeated three times and the average was tabulated.

Determination of ethanol
25 ml of asava were accurately measured and mixed with 100 ml of double distilled water and poured in to a separating funnel. The mixture was saturated with sodium chloride and added was 100 ml of hexane, shaken vigorously 2 – 3 min. The mixture was allowed to stand for half an hour. The lower layer was run in to a distillation flask. The hexane layer was washed with 25 ml of concentrated sodium chloride solution in a separating funnel then the NaCl layer was added to the distillation flask. The whole mixture was made alkaline with 1 M sodium hydroxide solution using solid phenolphthalein as indicator. To this added were a little pumice powder and 100 ml of water.

The whole mixture was distilled to obtain 90 ml of distillate. The distillate was poured in to a 100 ml volumetric flask and made the volume to 100 ml with double distilled water. Using this mixture relative density was determined to calculate the percentage v/v alcohol of the asava.

Determination of freezing point of asava
Freezing point is the maximum temperature occurring during the solidification of a super-cooled liquid. The apparatus for its determination was designed as that of the apparatus used in the determination of congealing point of asava.

About 5 ml of asava was placed in the inner test tube, which was immersed in a 500 ml capacited beaker containing water, fitted with a thermometer and a stirrer. The stirring was carried out at a rate of 25 cycles per min with simultaneous reduction in temperature by keep on adding ice cubes. When the temperature of the asava was observed to be 5°C or below, the beaker was filled with saturated NaCl solution to stabilize or maintain temperature. The process was continued until some seed crystals of asava were present. The process was repeated 3 times at least to get the average freezing point of asava.

Loss on Drying
About 10 ml (11.02 g) of the asava under study were accurately pipetted out and transferred to a tarred china dish which was known for its weight and kept in a hot air oven at 100 – 105°C for an hour. Then, the sample was weighed along with china dish to deduct the actual weight of tarred china dish. The weight of the content was noted to calculate the percentage loss on drying with reference to the asava.

Determination of loss of ignition
Though determination of loss on ignition is best suiting solid formulation like churna and the principle behind it is to convert all metallic oxalate, chloride, sulphate, phosphate, silicate et., in to their concerned oxide form.

Asava is a liquid formulation containing active principle in alcohol along with minerals in its aqueous layer or unfiltered fine crude drug particle during the preparatory moments. Hence, this
method of standardization was tried with 10 ml asava also using a silica crucible, after allowing asava be auto-evaporated at room temperature for about 1 h.

**Loss on Ignition**

A silica crucible was heated for about 30 min to red hot and cooled in a desiccator to note down its weight. About 10 ml of the asava was pipette out and then dried at 100 – 105°C for 1 h and ignited to constant weight in a muffle furnace at 600 - 625°C, until a carbon free ash formed. The crucible was allowed to cool in a desiccator after each ignition and care was taken to avoid catching fire. The weight of the carbon free ash was determined. The procedure was repeated to obtain a standard deviation to ensure consistency and then tabulated.

**Determination of pH of asava**

To determine the acidity or alkalinity of the asava at room temperature, potentiometric method was employed. The buffer solutions A – H were prepared using carbon dioxide free water as solvent as given in Indian Pharmacopoeia-1996 (A-95) which helped to detect the pH of asava whose range may be from 1.7 – 10.12.

**Determination of Refractive index** [22]

The refractive index (n) of a substance with reference to air is the ratio of the sine of angle of incidence to the sine of the angle of refraction of beam of passing from air in to the substance. The refractive index was conveniently measured using the Abbe refractometer at 25º C employing the wavelength of the D line of sodium (λ=589.3 nm), after calibrating the apparatus against distilled water whose nD^20 at 25º C was 1.3225.

**Determination of viscosity of asava**

The determination of viscosity of asava was carried out by means of capillary viscometer at room temperature. The viscometer was washed and dried completely. Then the viscometer was filled and examined through L tube to slightly above the mark G using a long pipette to minimize wetting the tube above the mark. The tube was placed vertically in a water bath maintained a temperature of 35º C and allowed to stand for half an hour to reach equilibrium. The volume of asava was adjusted so that the bottom of the meniscus settled at the mark G. The liquid was sucked to the point about 5 mm above the mark E and the pressure was revealed.

The time taken was measured for the bottom of the meniscus to fall from the top of mark E to the top edge of mark F. Then, the kinematic viscosity (V) in square mm per sec (mm^2 s^-1) using the expression

\[ V = K t \]

The constant (K) of the instrument was determined on a liquid of known viscosity (Dextran injection or saline).

**Determination of weight per ml of asava**

The weight per ml of a liquid is the weight, in g, of 1 ml of the liquid when weighed in air at room temperature. A thoroughly clean and dry Pycnometer was selected and filled with asava and weighed in air at room temperature. The procedure was repeated 3 times and average value of the weight of 1 ml of asava was calculated.
Primary organic analysis [23]
About 100 g of the crude drug (Patha) were powdered in a mechanical grinder, after a screening for the presence of foreign bodies, in to a moderately coarse powder were soxhleted successively with solvents of increasing polarity such as petroleum ether, benzene, chloroform and 75% ethanol (15 – 19 h) and a part of the extracts and the asava were subjected for the determination of and a primary organic analysis.

Primary organic analysis[24] of the both the extracts and the asava were carried out with suitable chemical reagents of research grade which led to a conclusion that the phenolic compounds were well pronounced.

Determination of total free sugar content in asava
The total free sugar content of asava was estimated using Benedict’s reagent for quantitative analysis and reported in terms of percentage w/ml as per the reference.

RESULTS AND DISCUSSION
The results of physical and physic-chemical analysis of asava were tabulated and discussed in detail under the section discussion (Table 1). The primary organic analysis on the both ethanolic extract of the crude drug (Patha) as well as the asava itself gave a positive test for carbohydrates (Molisch’s test); amino acid (Xanthoproteic test); free sugar (Fehling’s’ and Benedict’s test); tannins (Gold beater’s test); general phenolic compounds (dilute ferric chloride test); flavonoid (Shinoda’s test and pH dependent colour test by Mg-HCl); saponins (Haemolytic test); general glycosides (by hydrolytic test after exhausting free sugar); phenolic glycoside (by hydrolysis followed by phase separation by non-polar solvent and testing of the same); and the presence of anthocyanins (Blood red colouration of both alcoholic and aqueous extract) (Table 2). An organoleptic analysis was also carried out on the asava and the results were tabulated (Table 3).

Table 1. Standardization of asava by physical and physic-chemical methods

<table>
<thead>
<tr>
<th>S.No.</th>
<th>Parameters</th>
<th>Report/Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Total solids</td>
<td>47.8±0.25% w/ml</td>
</tr>
<tr>
<td>2.</td>
<td>Boiling range</td>
<td>79±0.06 – 109±0.08° C</td>
</tr>
<tr>
<td>3.</td>
<td>Congealing point</td>
<td>57±0.08 – 67±0.06° C</td>
</tr>
<tr>
<td>4.</td>
<td>Content of ethanol</td>
<td>21% v/v at 32° C</td>
</tr>
<tr>
<td>5.</td>
<td>Freezing point</td>
<td>11±0.06° C</td>
</tr>
<tr>
<td>6.</td>
<td>Loss on drying</td>
<td>19.82±0.50% w/w</td>
</tr>
<tr>
<td>7.</td>
<td>Loss on Ignition</td>
<td>2.8±0.55% w/v</td>
</tr>
<tr>
<td>8.</td>
<td>pH</td>
<td>4.4</td>
</tr>
<tr>
<td>9.</td>
<td>Refractive Index against water (1.332)</td>
<td>1.5777</td>
</tr>
<tr>
<td>10.</td>
<td>Viscosity against water (0.9982)</td>
<td>1.9987 poise at 32° C</td>
</tr>
<tr>
<td>11.</td>
<td>Weight per ml</td>
<td>1.166g/ml</td>
</tr>
<tr>
<td>12.</td>
<td>Total free sugar content</td>
<td>24.5 g % w/ml</td>
</tr>
<tr>
<td>13.</td>
<td>Fluorescence analysis (Long UV)</td>
<td></td>
</tr>
<tr>
<td>a.</td>
<td>Asava</td>
<td>brown</td>
</tr>
<tr>
<td>b.</td>
<td>Asava in water</td>
<td>Pale brown</td>
</tr>
<tr>
<td>c.</td>
<td>Asava with methanol</td>
<td>Yellowish brown</td>
</tr>
<tr>
<td>d.</td>
<td>Asava with ethylacetate</td>
<td>Pale brown</td>
</tr>
</tbody>
</table>

Results are presented as mean±Standard Deviation, n=3

It is noteworthy and deserves a mention here that the ethanolic extract of Viburnum secies has been proven to possess a remarkable antioxidant, anti-inflammatory and antiulcer activities. However, this drug, so far, has not been formulated in to any form and standardized for
its value. The asava itself and the asava added with water, 80% methanol and ethylacetate were observed under UV radiation showing brown, brown, yellowish brown and pale brown colouration respectively.

A primary organic analysis conducted on the asava itself as well as the ethanolic extract of the patha revealed the presence of carbohydrate, amino acid, free sugar, saponins, tannins, phenolic compounds (general), flavonoids, saponins and glycosides (phenolic glycosides). However, presence of phyto-sterols and triterpenes were in the negative. The asava was greenish brown in colour; aromatic in odour; aromatic and sweet in taste; sticky after minutes in texture between fingers; pourable and non-sticky in nature to view; it turned brownish green after its evaporation, when kept under room temperature; and smelled ethanolic and pleasant while heating on a boiling water bath.

Table 2. Primary organic analysis of asava against patha

<table>
<thead>
<tr>
<th>S.No.</th>
<th>Phytoconstituents</th>
<th>Asava</th>
<th>75% ethanolic extract of patha</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Carbohydrate</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>2.</td>
<td>Free sugar</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>3.</td>
<td>Amino acid</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>4.</td>
<td>Alkaloid</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>5.</td>
<td>Saponins</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>6.</td>
<td>Phyto-sterols</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>7.</td>
<td>Triterpenoids</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>8.</td>
<td>Tannins</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>9.</td>
<td>Flavonoids</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>10.</td>
<td>Glycosides (general)</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>11.</td>
<td>Glycoside (specific) (Phenolic glycosides)</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>12.</td>
<td>Anthocyanins</td>
<td>+++</td>
<td>+++</td>
</tr>
</tbody>
</table>

* - Test positive,  - Test negative

Table 3. Organoleptic analysis of asava

<table>
<thead>
<tr>
<th>S.No.</th>
<th>Parameters/Characters</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Colour</td>
<td>Greenish brown</td>
</tr>
<tr>
<td>2.</td>
<td>Odour</td>
<td>Aromatic</td>
</tr>
<tr>
<td>3.</td>
<td>Taste</td>
<td>Ethanollic and Sweet</td>
</tr>
<tr>
<td>4.</td>
<td>Texture</td>
<td>Sticky after minutes</td>
</tr>
<tr>
<td>5.</td>
<td>Nature</td>
<td>Pourable, Non-sticky</td>
</tr>
<tr>
<td>6.</td>
<td>Colour change at room temperature</td>
<td>darkening when volume reduced</td>
</tr>
<tr>
<td>7.</td>
<td>Odour upon heating</td>
<td>Ethanollic and pleasant</td>
</tr>
</tbody>
</table>

The term total solid is applied to the residue obtained where the prescribed amount of the preparation is dried to constant weight. The total solid of the asava were determined to be 47.8±0.25% w/ml. The lower limit of the range is the temperature indicated by the thermometer when the first drop of condensate leaves the tip of the condenser and the upper limit is the temperature at which the last drop evaporates from the lowest point in the distillation flask, as far as distilling range of the asava is concerned. In this event, the asava showed 79±0.06° C to 109±0.06° C as its boiling range.

The congealing temperature is that point at which there exists a mixture of the liquid (fused) phase of a substance and increasing proportion of solid phase. The asava, in this case, showed 57±0.08° C to 67±0.06° C as the congealing point. Making no modification in the setting of apparatus the freezing point of the asava was determined to be 11±0.06° C.
Since the principle behind the formulation of asava is that conversion of sugar (jaggery) into ethanol by anaerobic fermentation process, determination of total alcohol concentration was determined to be 21% v/v at 32°C by distillation cum specific gravity method. Loss on drying is a versatile method of standardization applicable for materials existing in liquid, solid, semisolid state. On the basis of the above principle, loss on drying of the asava was determined to be 19.82±0.60% w/w.

Although loss on ignition is best suiting to standardize formulation such as churna, it cannot be stated that asava may not be standardizable by this method. Because, the principle behind the loss on ignition is to determine the quantity of inorganic elements which could be convertible in to their corresponding oxides, which include both physiological as well as non-physiological ashes.

Hence, the loss on ignition of the asava in percentage w/v as determined to be 2.8±0.55% w/v. To determine the acidity or alkalinity of the asava, pH value was determined to be 4.4 by potentiometric method. Determination of refractive index is one of the best suiting standardizing process for liquid formulation with reference to air; the refractive index of the asava using as Abbe refractometer against water was measured to be 1.5777.

By employing an Oswald - type viscometer, viscosity was determined against water to be 1.9987 poise at 32°C. Since asava is a liquid formulation, by using a calibrated Pygnometer, the weight per ml of the asava was determined to be 1.166 g/ml at room temperature. The total free sugar content using Benedict’s reagent for quantitative analysis was determined to be 24.5 g %.

CONCLUSION

Every formulation of Ayurvedic system of medicine has its own modern scientific principle behind its preparation and standardization. The current study is to prove the same and to lay a path that further studies can be progressed on this phenolic rich formulation.

REFERENCES