Distribution, Nutritive Value and Mineral Composition of a Few Medicinal Plants of Shimoga District, Karnataka India

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Received: 25-05-2021; Revised: 28-07-2021; Accepted: 05-08-2021; Published on: 15-08-2021.

ABSTRACT
An attempt has been made to determine proximate, nutritive value, elemental composition and heavy metal contents of different flowers and fruits samples. The flowers and fruits samples were analyzed with the help of standard procedure. Among the macro elements nitrogen, potassium, calcium was dominant and phosphorus, magnesium was lowest in the respective fruit samples. The micronutrients study reveals that iron was the dominant element and manganese and copper was the lowest elements. Of the six proximates, moisture and carbohydrates was dominant and protein and fiber was the lowest in the different fruit samples. The highest value of nutritive value was recorded in the fruit sample of S. anacardium and lowest value of nutritive value was recorded in the fruit samples of B. frondosa. The average values of proximate, nutritive values were compared with recommended dietary allowances (RDA) values and discussed. Further, fruits and flowers help to development of ayurvedic formulation and to established cottage industries to improve economic and social condition of local people are appended.

Keywords: Butea frondosa Koen, Elemental composition, Nutritive value, Medicinal plants, Terminalia chabula Retz.

INTRODUCTION
Now-a-days animals and plants are suffering from numerous diseases, due to the lack of nutrition (Ash, moisture, fat, fiber, protein and carbohydrates) mineral (Na, K, Ca, Mg, P, Zn, Cu, Mn, Fe) and vitamins supplements. The specific deficiency of nutritional compounds, mineral components and vitamins causes the specific disorders like zinc deficiency causes pneumonia. Vitamin D deficiency causes osteoporosis, heart diseases, muscle and bone pain with undiagnosed causes, multiple sclerosis, rheumatoid, arthritis and various forms of cancer including breast, colon and prostate. Herbal drugs have been used by mankind since time immemorial to treat various disorders and offer an alternative to the synthetic compounds as they have been considered either non-toxic or less toxic. The traditional Indian system of medicine, ayurveda (Ayu-life, ved-knowledge) extensively uses the plant derived compound formulations for the treatment of various ailments after a careful study into the type of the disease. Plants are complex mixtures of compounds and no single compounds potentiate a desired therapeutic action while others reinforce the same and yet others interact to neutralize and counteract any possible side effects that may exist. Therefore, several plants with the common desired activities and varied undesirable activities are selected so that the final formulation will have a concentrated desired activity and the undesired activities will be diluted or absent altogether. Triphala is one of the ayurvedic formulation commonly prescribed by most health care practitioners in India. It is a traditional ayurvedic formulation consisting of the equal proportional mixture of dried fruits of three medicinal plants Amalaki (Emblica officinalis), Haritaki (Terminalia chebula) and Bibhitaki (Terminalia bellerica) and also known as ‘myrobalan’. Triphala means three (tri) fruits (phala). It is wild, non-habit forming and rejuvenates and hence is recommended for all. Tripala exhibits antiviral, antibacterial, antifungal and ant allergic properties and its constituents act as cardio tonic, control blood pressure, improve blood circulation and reduce cholesterol levels. Triphala shows immune modulator properties and helps in improving the body’s defense system. In recent years there are also several reports in the literature which suggest that triphala possess anti nut generic radio protecting and antioxidant activity.

T.chabula is belongs to family combretaceae it is used in fevers, cough, asthma, urinary diseases, piles, worms and rheumatism and scorpion sting. It is used in the preparation of triphala churna and has adjuncts to other medicines in numerous diseases like rheumatism. Terminalia bellerica fruits are useful in coughs, hoarseness, eye diseases and scorpion sting. It is used for preparation of Triphala churna, this churna is prescribed in diseases of the liver and gastro-intestinal tract and in a large variety of diseases. Kernel is narcotic and astringent and is used as an application to inflamed parts. Dried ripe fruit is astringent...
and employed in dropsy, piles and diarrhoea and also occasionally in fever. Gum is demulcent and purgative.\textsuperscript{16} 

\textit{Emblica officinalis} is belongs to family Euphorbiaceae. The fresh fruit is of the lungs and of the eves as a collyrium. In Persia it is used as vermifuge. The green fruits are made into pickles and preserves to stimulate appetite. This is a nutritive tonic useful in phthisis, and improves all conditions of debility. The drug is also used in scorpion-sting.\textsuperscript{16} 

\textit{Sapindus trifoliatus} is belongs to family Sapindaceae. The seed of \textit{Sapindus trifoliatus} is used to stimulate the uterus to child birth and in amenorrhea. Seeds pounded up with water and introduced into the mouth cut short the paroxysm of epilepsy.

\textit{Semicarpus anacardium} is belongs to the family Anacardace. The tree is large handsome with aspanicles of small yellow male flower and inconspicuous greenish females. Wood is soft grayish white, and useful. Fruit it is used for strength of teeth. The seed oil is specific in rheumatism.

\textit{Butea frondosa} is belongs to the family Fabaceae. Bark furnishes a very important exudation which hardens into a red brittle resin known as butea gum or Bengal kino. Medicinally it is an excellent astringent, useful in diarrhoea and dysentery. Fresh juice is also applied to ulcers and relaxed. Congested and septic sore throat; internally it is given in diarrhoea dysentery and phthisis. As anthelmintic and aperients. Flowers are used for dyes. Seeds are useful in ring worn, herpes, bark is given with ginger in snake bites. A week decoction of the bark is useful in catarrh, cold and cough.\textsuperscript{16} In the present study, medicinally important plants of \textit{Terminalia chebula}, \textit{Terminalia bellerica}, \textit{Emblica officinalis Sapindus trifoliatus}, \textit{Semicarpus anacardium} and \textit{Butea frondosa} were taken for investigation from the different places of Shimoga district.\textsuperscript{17} It is the three constituents that is Amalaki, Bibhitaki and Harataki represents these three humors Amalaki is related with pitta humor, so it helps treating inflammatory complications, liver problems, ulcers, constipation, diarrhea, infections and many others. Apart from that, Amalaki offers its significant role as antibacterial and antiviral substance, pronounced expectorant and cardiovascular nourishment tonic. Bibhitaki corresponds to kapha humor. It helps treating asthma, allergies, coughs and bronchial complications. Haritaki is linked with vata humor. It is helpful in treating chronic constipation of these three constituents.\textsuperscript{18} All of these constituents have been studied scientifically and the result confirms the traditional benefits of the same. In total, it is one of the best ayurvedic rasayana or preparation that others an excellent solution for lowering cholesterol level, reducing high blood pressure, enhancing blood circulation, improving digestive system and managing eradication without being dependent on laxatives. Each of the fruit components of Triphala takes a great care of the body system by promoting gentle purification of all toxic elements of the body, while improving the digestive system throughout. Due to its ability to nourish internal organs of the body. It resembles the care of the mother to her child. With its high nutritional value, it cleanses the body at the deepest organic level without running down the reserves of the body system. That makes the preparation one of the best among all herbal mediations in the world.\textsuperscript{15} It exhibits antiviral, antibacterial, antifungal and antiinflammatory properties.\textsuperscript{16-17} Triphala and its constituents act as cardio tonic, control blood pressure, improve blood circulation and reduce cholesterol levels. Triphala shows immune modulators properties and helps in improving the body’s defense system. In recent years there are also several reports in the literature which suggest that triphala possess ant mutagenic radio protecting and antioxidant activity.\textsuperscript{18-19} Kumar \textit{et al.} showed that radio protective effects of Rasayanas.\textsuperscript{20} Rama Sundaran Srikumar \textit{et al.}, studied effect of Triphala an oxidative stress and on cell-mediated immune response against noise stress in rats.\textsuperscript{21} Recently, Naik \textit{et al.},\textsuperscript{22} studied free radical scavenging reactions and phytochemical analysis of Triphala, an ayurvedic formulation and Rukmini and Udaya shankara Rao studied the chemical and nutritional values of kernel and its oil of \textit{Terminalia bellerica} Roxb. is a valuable tree of Indian forests.\textsuperscript{23} However, with the nutritional values and elemental composition of fruits of components of Triphala.\textsuperscript{24} In the present study, medicinally important plants of \textit{Terminalia chebula}, \textit{Terminalia bellerica}, \textit{Emblica officinalis Sapindus trifoliatus}, \textit{Semicarpus anacardium} and \textit{Butea frondosa} were taken for investigation from the different places of Shimoga district.

\textbf{MATERIALS AND METHODS}

\textbf{Study Area}

Shimoga district lies towards the North-western part of Karnataka state. It is situated between 13°27’to 14°39’ North latitude and 74°38’ to 76°4’ East longitude. The district stretches 154 kms East-West direction and 128 km in North South direction. The district is bordered by Davangere district in the East, Chikmagalur in the south and Dharwad district in the North. On the North-West border, North Canara and on the south-west by South Canara (Map 1).
The main rivers of the district are the Tunga, Bhadra and Sharavathi. The river Tunga runs in most part of the district. It joins with Bhadra at Kudli. The Sharavathi flows towards North-West taking its origin near Kavale Durga. It hurls down the Ghat at Jog. Creating the world famous Jog falls. The Gargita river originates a Kodachadri and flows downward towards South Canara through Haidar Ghar Ghat. There are several tanks in the district of. Of these, the Shantisagar tank is the largest. As greater part of the district consists of several mountains covered by thick forest, the climate is very agreeable. The average rainfall of the district is 827 cm towards western side and 90 cm towards the Eastern boundary. The rainy season extend from May-June to October-November, December-April practically there is no rain. The hottest period is between March to May (http://en.wikipedia.org.Karnataka).  

Collection of plants and distribution

The medicinal plants were collected and distribution study was made on the basis of regular field visits. The plants were collected and materials were brought to the laboratory and technical descriptions were made on the basis of observations. The plant were identified and assigned them to the respective taxonomy position. Simultaneously, various taxonomical literatures were also referred to confirm their distribution and identity.

Medicinal values

The medicinal uses of identified plants were documented on the basis of literature.

Preparation of plant sample for analysis

The collected plants were carried to the laboratory as quickly as possible using polythene bags. The samples were washed with the help of camel hairbrush and sponging with a piece of cotton wool in a 0.1% detergent solution or dilute (0.1% N) HCl to remove the adherent dust particles, then it was rinsed into lots of distilled water and remove the excess of water from the plant surface using blotting paper or ordinary filter paper and separated the flowers and fruits portions for separate analysis, cut the plant sample into pieces of suitable size. After cleaning, the sample was shade dried in a dust fume free place, not sun light because nitrogen fractions may be lost through volatilization. Then material was shade dried it was grinded to fine powder using pestle and mortar to avoid contamination (Plate-1). Then powder was preserve in an air tight bottles (jars) or pressing covers and store the material under the refrigerated condition for further analysis of nutrients.

Determination of Nutritive Value

Determination of Ash content

10 g of each sample was weighed in a silica crucible. The crucible was heated first over a low flame till all the material was completely charred, followed by heating in a muffle furnace for about 3 to 5 h at 600°C. It was cooled in a desiccator and weighed to ensure completion of ashing. It was heated again in the furnace for half an hour, cooled and weighed. This was repeated constantly till the weight become constant (ash become white or grayish white) Weight of the ash giving the ash content.  

\[
\text{Ash } \% = \frac{\text{Loss of weight of sample}}{\text{Initial weight of sample}} \times 100
\]

Determination of moisture content

The sample materials were taken in a flat bottom dish and kept overnight in an air oven at 100-110°C and weighed. The loss in weight was regarded as a measure of moisture content (NISCOM, CSIR, 1999). The percent of moisture has been calculated using the following formula.

\[
\text{Moist content } \% = \frac{\text{Loss of weight of sample}}{\text{Initial weight of plant sample}} \times 100
\]

Determination of crude fat

The crude fat determined by extracting 2 gm of moisture free sampling with petroleum ether in a Soxhlet extractor, heating the flask on a sand-bath for about 6 h till a drop taken from the drippings left no greasy stain on the filter paper. After boiling with petroleum ether (Chopra and Kanwar, 1991) and the residual petroleum ether was filtered using Whatman No. 40 filter paper and the filtrate was evaporated in a pre-heated beaker. Increase in weight of beaker gave the crude fat. The percent of crude fat has been calculated by using the following formula.

\[
\text{Crude fat } \% = \frac{\text{Weight of the crude fat}}{\text{Weight of the plant material}} \times 100
\]

Determination of crude fiber

The crude fibre was determined by titrating the moisture free and fat free materials were treated with 200 ml of 1.25% sulfuric acid. After filtration and washing, the residue was treated with 1.25% of sodium hydroxide solution. It was filtered, washed with hot water and then 1% HNO3 and again with hot water. The residue was ignited and the ash was weighed. The loss in the weight gave the weight of the crude fibre (Chopra and Kanwar, 1991). The percent of crude fibre can be calculated by using the following formula.

\[
\text{Crude fibre } \% = \frac{\text{Weight of the ash}}{\text{Weight of the plant material}} \times 100
\]

Protein Estimation by Lowry's method

Sample preparation

Extraction was carried out with buffer used for the enzymes assay. 500mg of sample was taken and grind by using pestle and mortar in 5-10 ml of the buffer, then centrifuged and used the supernatant for protein estimation.
Estimation of protein

Pipette out 0.2, 0.4, 0.6, 0.8 and 1ml of working standard solution was taken in a series of test tube. Then 0.1 and 0.2 ml of plant sample was pipetted out in two other test tubes and make up the volume to 1 ml with distilled water taken as blank. Then added 5 ml of regent C to each tube including the blank and mixed well and allowed to stand for 10 min. Then added 0.5 ml of reagent D again mixed well and incubated at room temperature in the dark for 30 min then blue colour was developed. The reading was taken at 660 nm and a standard graph was drawn and calculated the amount of protein in sample.33-37

Determination of carbohydrate

The percent of carbohydrate was calculated by using the following formula as follows,

Carbohydrate % = 100 – [ash % + moist % + fat % + protein %]

Determination of nutritive value

The percent of nutritive value was calculated by using the following formula as follows,

Nutritive value % = 4 x protein % + 9 x fat % + 4 x carbohydrate %

Preparation of plant Sample for mineral Analysis

One gram of powdered dried plant material was taken in 250 ml of conical flask 25 to 35 ml of diacid mixture (900 ml HNO₃ + 400 ml perchloric acid) was added, each conical flask was closed by using funnels. Then digestion was carried out by using hot plates at 1 h up to get clear solution at 10 minutes then added 50 ml of distilled water and filtered by using Whatman filter paper. After completion of filtration, the content was transferred quantitatively to 100 ml of volumetric flask and final volume was adjusted to 100 ml by adding distilled water. This solution was ready for mineral analysis.33-37

Determination of Macronutrients

Determination of sodium and potassium

Principle of flame photometry

The aliquot of plant digest containing alkali and alkaline earth metallic cations is fed to the flame photometry, it vaporizes to gaseous state. The compound dissociates into individual atoms which get excited due to the heat energy of flame. On the excited the electrons jumps from its normal orbit to higher energy level. The excitation state of the atom does not remain for longer time as it is unstable, it reverts to its original level. While returning to its original state, the atom loses the energy in the form of radiation. The characteristic colour and wave length of the radiation indicates the type of the element. Yellow coloured radiation with 589 nm wave length indicates presence of sodium and red coloured radiation with wave length 766 nm indicates the presence of potassium.33-37

Materials and reagents required

1) 250 ml beaker with glass rod.
2) 100 ml volumetric flask.
3) Pipettes 1, 2, 5 and 10 ml.
4) 1 litre volumetric flask.
5) 50 ml volumetric flask.
6) Electric or chemical balance

Preparation of sodium standards

100 ppm sodium stock solution was prepared by dissolving 0.6304 gm of pure dried sodium chloride in distilled water. 100 ppm sodium stock solution was prepared by diluting 10 ml of 1000 ppm sodium stock solution. The various working sodium standard solution of 0, 1, 2, 3, 4 and 5 ppm were prepared with suitable dilution.

Preparation of potassium standards

1000 ppm sodium stock solution was prepared by dissolving 1.908 gm of pure dried KCl in distilled water. 100 ppm sodium stock solution was prepared by diluting 10 ml of 1000 ppm sodium stock solution. The various working sodium standard solution of 0, 1, 2, 3, 4 and 5 ppm were prepared with suitable dilution.

Procedure

2 ml of aliquot plant digested material was taken in 50 ml volumetric flask and make up the volume adding distilled water. This gives a dilution factor of 50. Then adjusting the flame photometer reading to ‘0’ with ‘0’ ppm standard and 100 with 5 ppm standard solution and flame photometer reading for other standard feeding the diluted solution of plant digested and recorded the flame photometer. Reading and draw the standard curve of sodium / potassium by plotting the flame photometer readings along with the y-axis and concentration along with the x-axis, detect the concentration of Na / K in the curve.33-37

The % of Na / K was calculated with the help of following formula.

\[
\% \text{ of Na}/K = \frac{\text{Graph ppm}}{10^6} \times \text{Dilution factor} \times \frac{\text{Volume of plant digestion made}}{\text{Weight of the plant sample}} \times 100
\]

Orthophosphate (phosphorus) present in the plant digest when react with vanadate and molybdate gives a yellow coloured complex i.e. phospho-vanadomolybdate in acid solution. The yellow colour due to the substitution of
oxyvanadium and oxyxymolybdenum radicals for oxygen of PO₄ to give a heteropoly compound. The intensity of yellow colour is measured colourimetrically at 400 to 490 nm using spectrophotometer.

**Reagents required**

Vanadomolybdate reagent (Ammonium molybdate-Ammonium vanadate in nitric acid).

**Preparation of solution A**

Dissolve 1.25 g ammonium metavanadate in 300 ml of boiling water. Cool and add 250 ml of concentrated HNO₃ and again cool to room temperature.

**Preparation of solution B**

Dissolve 25 g of ammonium molybdate in 400 ml of distilled water pour solution A to solution B mix well and make up the volume to 1 litre with distilled water.

**Preparation of phosphorus standards**

**Preparation**

Prepare 100 ppm of P standard stock solution by dissolving 0.2195 g of pure KH₂PO₄ in 500 ml of distilled water.

\[
% \text{ of } P = \left( \frac{\text{Graph ppm}}{10^6} \times \frac{\text{Volume of dilution made}}{\text{Aliquot}} \right) \times \frac{\text{Volume of plant digest}}{\text{Weight of plant sample}} \times 100
\]

**Determination of calcium and magnesium**

**Procedure**

1 ml of aliquot plant digested material was taken in 50 ml of volumetric flask and 10 ml of vanadomolybdate reagent was added and mixed thoroughly and final volume was adjusted to 50 ml by adding distilled water. Feed the sample to AAS and observe the results with the help of calcium / magnesium hallow cathode lamp with a wave length 422.7 and 28.2 nm of calcium / magnesium. Draw the calibration curve with the help of concentration value.

\[
% \text{ of } \text{Ca} / \text{Mg} = \left( \frac{\text{Graph ppm}}{10^6} \times \text{Diution factor} \right) \times \frac{\text{Volume of plant digestion made}}{\text{Weight of the plant sample}} \times 100
\]

**Determination of Nitrogen**

**Principle**

Nitrogen content of plant was converted into (NH₄)₂SO₄ by digesting with diacid mixture (concentrated H₂SO₄). The acid digest was distilled for ammonia by using 40% NaOH. The distilled ammonia was trapped in boricacid mixed indicator solution. The amount of ammonia trapped was estimated by titrating against standard acid.

**Reagents**

1) Concentrated H₂SO₄ or diacid mixture (9 : 4 = HNO₃:HClO₄(Perchloric acid )
2) 40 % NaOH
3) 2% or 4% boric acid
4) 0.01 N H₂SO₄
5) Mixed indicator (Bromeresol green) (0.1g) + Methyle red (0.07g) in 100 ml of 95 % of ethanol.

**Working standards of phosphorous**

Transfer 0, 1, 2, 2.5, 5, 7.5 and 10 ml of 100 ppm P-stock solution into separate 50 ml volumetric flask to get 0, 2, 5, 10, 15 and 20 ppm of P-working standards.

**Procedure**

Transfer 5 ml of aliquot of plant digest (Triacid or diacid digested plant samples) into 50 ml volumetric flask. Add 10 ml of vanadomolybdate reagent to samples and also to each standards and mix thoroughly and make up the volume to 50 ml with distilled water. After 30 minutes of colour development read the intensity of yellow colour on a spectrophotometer at 470 nm (i.e., between 400-490 nm). Draw the calibration curve (standard graph) of P standards by plotting the P-absorbance against P-concentration.

1. Find out the P-content in plant digest sample by referring to the standard curve.

**Calculation**

The % of P in the plant sample was calculated with a help of following formula.

\[
% \text{ of } P = \left( \frac{\text{Graph ppm}}{10^6} \times \frac{\text{Volume of plant digest}}{\text{Weight of plant sample}} \right) \times 100
\]
10 ml of diluted acid digested sample was taken in a micro Kjeldal distillation assembly. Boric acid mixed indicator solution was kept at ready at the receiving end to trap ammonia, 30 ml of 40% NaOH was added and distillation was carried out till the colour of the mixture changes from bluish purple to bluish green further the distillation is also continued to for some of time to trap all the ammonia, released. The completion of distillation was confirmed by using litmus test in which no changes in colour of the red litmus paper indicate the complete distillation. The quantity of ammonia distilled was estimated by titrating against 0.01 N H₂SO₄ or HCl till the colour changes to purple. 33-37 The % of N was calculated with the help of following formula.

\[
\text{% of Nitrogen} = \frac{\text{Titrate value} \times N \text{ H}_2\text{SO}_4 \times 0.014 \times \text{dil factor (if any)}}{\text{Weight of plant sample (g)}} \times 100
\]

### Determination of Micronutrient cations in plant sample by using Atomic Absorption Spectrophotometry (AAS)

Micronutrients namely Zn, Fe, Cu and Mn were analyzed from the plant samples with the help of AAS. The digested samples which were prepared for the elemental analysis are used with the following modifications.

The 2 ml of digested samples were taken and diluted to 50 ml and the following micronutrients are analyzed.

\[
\text{ppm of micronutrients} = \frac{\text{ppm of micronutrients in plant}}{1000} \times \frac{\text{Vol. of plant digest}}{\text{Weight of the plant sample}} \times \text{dilution factor} \times 1000
\]

### Analysis of Cadmium and Lead

In addition to micro and macro-elements, cadmium and lead were also analyzed. The procedures and calculations are followed as described in the micro elemental analysis. 33-37

### RESULTS AND DISCUSSION

Of the six medicinal plants which were studied for distribution, mineral composition and nutritive value was discussed (Table 1 and 2).

**Terminalia chebula** Retz. (Fig 5.A.) Ramaswamy et al., 27 2001 reported the four species of **Terminalia** form Shimoga district. The plant is reported from Ullur of Sagar taluk, where plant is scatteredly found in association with *Xyilia xylocarpa* (Roxb.) Taub. and *Holarrhena antidysenterica* (Roth.) DC however, the plant is very rare in the Kuvempu University campus. It is also found in Srilanka, Indomalaya, Burma and Kampuchea. **Terminalia bellerica** Roxb (Fig 5.B) is recorded in the deciduous forests of all districts of Madras presidency up to the altitude of 3,000 ft. Ramaswamy et al., 27 2001 recorded **Terminalia bellerica** in the forest of Kattinakare of Sorab taluk, Ullur of Sagar taluk and Gajanur of Shimoga taluk of Shimoga district and they have observed in the distribution **Terminalia bellerica** often in association with **Terminalia crenulata** (Roxb.) Roth. and *Ervatamia heyneana* (Wall.) Cooke. The plant is frequently found in the Kuvempu University campus which belongs to Bhadra Wildlife Sanctuary. Sri Rambhat (1992-1993) 28 recorded **Terminalia bellerica** from Kodachadri forest of Shimoga district. Gamble (1998) 26 classify species of **Emblica officinalis** Goertn (Fig 5.C.) and **Emblica fischer** (Gamble, 1998) 26.

### Table 1: Distribution of medicinal plants in various taluks of Shimoga district

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Name of the plants</th>
<th>Taluk</th>
<th>Sagar</th>
<th>Sorab</th>
<th>Shikaripura</th>
<th>Thirthahalli</th>
<th>Bhadravathi</th>
<th>Shimoga</th>
<th>Hosanagara</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><strong>Terminalia chebula</strong></td>
<td></td>
<td>+</td>
<td>+</td>
<td>No</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td><strong>Terminalia bellerica</strong></td>
<td></td>
<td>+</td>
<td>+</td>
<td>No</td>
<td>+</td>
<td>+</td>
<td>No</td>
<td>+</td>
</tr>
<tr>
<td>3</td>
<td><strong>Emblica officinalis</strong></td>
<td></td>
<td>+</td>
<td>+</td>
<td>No</td>
<td>+</td>
<td>+</td>
<td>No</td>
<td>+</td>
</tr>
<tr>
<td>4</td>
<td><strong>Sapindus trifoliatus</strong></td>
<td></td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>5</td>
<td><strong>Semicarpus anacardium</strong></td>
<td></td>
<td>+</td>
<td>(Banajalaya)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>6</td>
<td><strong>Butea Frondosa</strong></td>
<td></td>
<td>+</td>
<td>+</td>
<td>No</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

Plants observed; ‘+’ Not seen; ‘No’ Not observed.
The species are classified on the basis of length of the branchlets shape of the fruits and the floral characteristic features. The *Emblica officinalis* was recorded in the North Circars, Deccan and Carnatic, where it was found in the dry deciduous forests of up to the altitude of 4,000ft. The species is predominantly found in Mahendragiri and dry slopes of Western Ghats. It is important note that, though Gamble (1998) referred *Emblica fischeri* (Gamble, 1998) in the flora of Madras presidency, he referred that he did not observed the fruits. Ramaswamy *et al.*,27 mentioned three species of Phyllanthus Linn. *Emblica officinalis* Gaertn observed in the deciduous forest of Humcha taluk of Shimoga district. The plant was scatteringly distributed. A few plants were observed in the premises of ladies hostel, top the Shankarghatta and behind the employment office of the Kuvempu University campus. Venugopal (1992-1993)29 recorded *Phyllanthus emblica* Linn. at Jade of Sorab taluk of Shimoga district. *Butea frondosa* Koen (Fig.5 E) (Gamble (1998) recorded to species of *Butea frondosa* Koen and *Butea superba* Roxb was observed in the open country and deciduous forest perits of Deccan. The plants were common on black cotton soil and salts lands. The plant yields a red kinogum and lack insect is grown upon it. *Butea superba* Roxb, the leaves and flowers and pods were like those of *Butea frondosa*. However, the flower rather more yellow slight large than that of frondosa. The plant also yields kinogum and flowers yield red dye. Ramaswamy *et al.* (2001)21 reported *Butea monosperma* Lan, *Butea frondosa* Koenig ex Roxb. and *Plano monosperma* Lam Kuntz. It was found in the Sampagodkan forest of Sorab taluk. Jog of Sagar taluk of Shimoga district. Sri Rambhat (1994)28 reported *Butea monosperma* from Kodachadri forest and said that the plant is used as astringent. *Sapindus trifoliatus* Linn (Fig 5 D.) Gamble (1998) recorded two species of *Sapindus* Linn. *Sapindus lourifolia* vahal; which was recorded from Western Ghats of South canara and Mysore to the Anamalais and hills of Madura where it was found in the evergreen and open forests at low

### Table 2: Proximates, nutritive value, and mineral components of medicinal plants of Shimoga district

<table>
<thead>
<tr>
<th>Sl No</th>
<th>Factors</th>
<th><em>T.chabula</em></th>
<th><em>T. bellerica</em></th>
<th><em>E.officinalis</em></th>
<th><em>S.trifoliatus</em></th>
<th><em>S anacardium</em></th>
<th><em>B.frondosa</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Macronutrients</td>
<td>1.56±0.18</td>
<td>2.34±0.21</td>
<td>3.13±0.88</td>
<td>1.71±0.11</td>
<td>1.81±0.10</td>
<td>1.8±0.61</td>
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<td>Na</td>
<td>4.04±0.05</td>
<td>3.37±0.11</td>
<td>3.07±0.06</td>
<td>3.50±0.24</td>
<td>1.73±0.20</td>
<td>5.69±1.89</td>
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<td>K</td>
<td>0.30±0.005</td>
<td>0.34±0.018</td>
<td>0.23±0.003</td>
<td>0.69±0.017</td>
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<td>0.52±0.22</td>
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<td>0.41±0.02</td>
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<td>Mg</td>
<td>3.55±0.05</td>
<td>3.40±0.06</td>
<td>4.36±0.02</td>
<td>8.82±0.00</td>
<td>2.17±0.02</td>
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<td>II</td>
<td>Micronutrients</td>
<td>88.00±14.14</td>
<td>50.83±1.32</td>
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<td>89.71±6.22</td>
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<td>Zn</td>
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<td>65.06±4.41</td>
<td>38.73±11.75</td>
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<td>Cu</td>
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<tr>
<td>IV</td>
<td>Nutritive value</td>
<td>10.27±1.30</td>
<td>10.90±0.55</td>
<td>4.50±0.50</td>
<td>11.80±1.68</td>
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<td>167.40±0.50</td>
<td>83.20±4.60</td>
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<td>108.00±12.17</td>
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<td>96.50±17.90</td>
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<td>Fibre%</td>
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<td>Protein%</td>
<td>9.34±3.77</td>
<td>20.26±0.96</td>
<td>150.04±9.17</td>
<td>100.25±7.39</td>
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<td>53.39±7.50</td>
</tr>
<tr>
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<td>Carbohydrates %</td>
<td>940±1.69</td>
<td>958.34±1.62</td>
<td>882.90±2.93</td>
<td>1174.08±7.36</td>
<td>1117.08±7.36</td>
<td>561.90±5.38</td>
</tr>
</tbody>
</table>

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elevations he mentioned that the fruit is used like that of the soap. Pasakotta is the vernacular name of Malayalam. *Sapindus emarginatus* is observed North Circars, Deccan and Carnatic, extending into Eastern slopes of the Nilgriris and Pulneys and the hills of Tinnevelly of Madras presidency. The tree was frequently found in the deciduous and dry evergreen forests, frequent on the Coast of Sriharikota in Nellore; it is also cultivated. The fruit is used for washing and commonly known as soap nut. The plant is observed between Kargal and Jog by Ramaswamy *et al.*, 2001. The plant is planted in Kuvempu University Campus as avenue tree along the margin of the road side. Venugopal (1992-193) recorded from the village Sanda. *Semiarpus anacardium* Linn. Three species of *Semicarpus anacardium* Linn. were recorded from the Madras presidency (Gamble 1998) *Semicarpus anacardium* Linn. A moderate sized tree whose pericarp of the drupe gives a marking ink and hypocarp is edible. *Semicarpus travancorica* Bedd which was recorded in the Western Ghats of evergreen forests of Travancore and Tinnevelly up to the altitude of 4,000ft. The plant is a large tree with grayish-white wood which has no value. *Semicarpus anacardium* was also observed in the evergreen forest of Travancore and Tinnevelly upto the altitude of 4,000ft. The tree is large handsome with a panicles of small yellow male flowers and inconspicuous greenish females. Wood is soft grayish-white, and useful. *Semicarpus anacardium* the mark nut tree was found in the deciduous forests of all district of Madras Presidency. Ramaswamy *et al.*, 2001 recorded the plant at Bilesvarra, Kattinakere, between Hosanagara and Sagar taluk of Shimoga district. The plant is scatteredly distributed in the deciduous forests. Venugopal (1992-1993) recorded *Semicarpus anacardium* Linn. at Marsa of sagar taluk and mentioned that the receptacle of fruit and seed oil and fruits are medicinally used (Map 1 and Table 1).

Macronutrients

Of the six macronutrients, the highest percentage value of 3.13 ± 0.88 sodium was recorded in the fruit sample of *E.officinalis* and lowest percentage value of 1.56 ± 0.18 was recorded in the fruit sample of *T.chabula*, whereas in case of potassium the highest percentage value of 5.69 ± 1.89 potassium was recorded in the flower sample of *B.frondosa* and lowest percentage value of 1.73 ± 0.20 potassium was recorded in the fruit sample of *S.anacardium*. The highest percentage value of 0.69 ± 0.017phosphorus was recorded in the fruit sample of *S.trifoliatus* and lowest value of 0.23 ± 0.003 phosphorus was recorded in the fruit sample *E.officinalis*. The highest value of 6.20 ± 3.16 calcium was recorded in the fruit sample of *S.anacardium* and lowest value of 0.52 ± 0.22 calcium was recorded in the fruits of *S.trifoliatus*. The highest percentage value of 0.87 ± 0.02 Magnesium was recorded in the fruit sample *B.frondosa* and lowest value of 0.20 ± 0.02 was recorded in the fruit sample *E.officinalis*. The highest percentage value of 8.82 ± 0.00
Nitrogen was recorded in the fruit sample of *S. trifoliatus* and lowest value of 2.17 ± 0.02 was recorded in the fruit sample of *S. anacardium* (Table 2 and Fig.1).

Figure 1: Variation of macro nutrients (%) in different medicinal plants

*T.chabula*: Potassium > Nitrogen > Sodium > Calcium > Phosphrous > Magnesium

*T.bellerica*:
Calcium>Nitrogen>Potassium>Sodium>Magnesium>Phosphrous

*E.officinalis*:
Sodium>Calcium>Nitrogen>Potassium>Phosphrous>Magnesium

*S.anacardium*:
Calcium>Nitrogen>Potassium>Sodium>Magnesium>Phosphrous

*B frondosa*:
Potassium>Nitrogen>Potassium>Calcium>Phosphrous>Magnesium

**T. chabula** Iron>Zinc>Copper>Manganese

**T. bellerica** Iron>Copper>Zinc>Manganese

**E. officinalis** Iron>Zinc>Manganese>Copper

**S. trifoliatus** Iron>Zinc>Manganese>Copper

**S. anacardium** Iron>Manganese>Copper

**B. frondosa** Manganese>Copper>Zinc>Iron

Of the four micronutrients the highest ppm of 204.86 ± 9.96 zinc was recorded in the fruit sample of *S. trifoliatus* and lowest ppm of 40.93 ± 1.12 zinc was recorded in the fruit sample of *B. frondosa*. The highest ppm of 89.97 ± 30.15 copper was recorded in the fruit sample of *B. frondosa* and lowest ppm of 29.24 ± 3.61 copper was recorded in the fruit sample of *S. anacardium*. The highest ppm of 680.03 ± 228.84 manganese was recorded in the fruit sample of *B. frondosa* and lowest ppm of 23.41 ± 1.52 manganese was recorded in the fruit sample of *T.chabula*. The highest ppm of 910.48 ± 331.43 iron was recorded in the fruit sample of *S. anacardium* and lowest ppm of 19.5 ± 6.5 iron was recorded in the fruit sample of *B. frondosa* (Table 2 and Fig 2).

Figure 2: Variation of micro nutrients (ppm) in different medicinal plants

**Proximates and Nutritive value**

*T.chabula*
Moisture>Fat>Ash>Protein>Carbohydrates>protein

*T.bellerica*
moisture>Fat>Protein>Carbohydrates>Ash>Fibre

*E.officinalis*
Carbohydrates>Moisture>Ash>Fat>Fibre>Protein

*S. trifoliatus*
Carbohydrates>Fat>Moisture>Ash>Protein>Fibre

*S. anacardium*
Moisture>Carbohydrates>Ash>Fat>Protein>Fibre

*B.frondosa*
Moisture>Carbohydrates>Fat>Ash>Fibre>Protein

Nutritive value *S.anacardium>*S trifoliatus>*T.bellerica>*E.officinalis>*B.frondosa*

The highest value of ash 19.50 ± 0.50 was recorded in the fruit sample of *B.frondosa* and lowest value of 10.27±1.30 ash was recorded in the fruit sample of *T.chabula*. The highest value of 185.00 ± 7.64 moisture was recorded in the fruit sample of *B.frondosa* and lowest value of 83.20 ±4.60 moisture was recorded in the fruit sample of *E.officinalis*. The maximum value of 103.50±7.63 fat was recorded in the fruit sample of *T.chabula* and minimum value of 4.50±0.26 fat was
The result of the macronutrients, micronutrients, proximate and nutritive value is compared with recommended dilatory allowance (RDA) (Table 3).

Indrayana et al. (2005)\textsuperscript{11} said that Na and K take part in ionic balance of the human body and maintain tissue excitability. Because of the solubility of salt, Na plays an important role in the transport of metabolites. K is of important as a diuretic. Calcium constitutes a large proportion of the bone, human blood and extracellular fluid. It is necessary for the normal functioning of cardiac muscles, blood coagulation and milk clotting and the regulation of cell permeability. It also plays an important part in nerve impulse transmission and in the mechanism of neuromuscular system. Mg is required in the plasma and extracellular fluid where it helps maintain osmotic equilibrium. It is required in many enzyme catalyzed reactions especially those in which nucleotide participate where the reactive species in the magnesium salt example, MgATP2, lack of Mg is associated with abnormal irritability of muscle and convulsions and excess of mg with depression of the central nervous system. Further, Indrayana et al. (2005)\textsuperscript{11} discussed the importance of sodium, potassium, calcium and magnesium of medicinally valued plants of Uttarakanchal. Trace metals composition of food is of interest because of their essential toxic nature (Onianwa et al., 1999).\textsuperscript{38} Micronutrients constitute a small fraction of the entire diet but play important role in different metabolic processes (Akhter et al., 2002).\textsuperscript{39} Wide variations in concentrations of trace metals have been reported in bovillon cubes, mixed species and nuts (Akpasyung, 2005; Garcia et al., 2005; Satter et al., 1989 and Ansari et al., 2004).\textsuperscript{40-43}

Micronutrients like namely Zn, Cu, Mn and Fe were analyzed in the parts which have been been used as of Terminalia chebula, Terminalia bellerica and Emblica officinalis. Mixture of medicinal plants are prescribed by the traditional healers for diseases ranging from common cold to malaria, arthritis, ulcers etc (Obaijunwa et al., 2002).\textsuperscript{44}

The cadmium and lead was not recorded in the plant parts which are studied except in the fruit samples of Terminalia chebula and Terminalia bellerica in which traces of cadmium and lead was recorded. Lead and cadmium along with iron, Mn, Cu, Ni, Zn etc., are also considered as heavy metals. Al Mourut Alukayode Ajasa et al. (2003)\textsuperscript{45} studied heavy metals and macronutrient status in herbal plants of Nigeria (2004). Obaijunwa et al. (2002) studied essential and trace metals constituents of some Nigerian medicinal plants and reported that certain toxic elements such as lead, arsenic, mercury, cadmium etc., which are prime intact in toxicological study (War et al., 1996) were not found from the plant samples which they have studied.

recorded in the fruit sample \textit{E.officinalis}. The maximum value of 13.28±5.78 fibre was recorded in the fruit sample of \textit{B.frondosa} and minimum value of 3.00±0.00 fibre was recorded in the fruit sample of \textit{S.trifoliatus}. The highest value of 8.38±0.22 of protein was recorded in the fruit sample of \textit{T.chabula} and lowest value of 1.44 ± 0.06 protein was recorded in the fruit sample of \textit{B.frondosa}. The maximum value of 150.04 ± 9.17 carbohydrates was recorded in the fruit sample of \textit{E.officinalis} and minimum value of 20.26 ± 0.96 carbohydrates was recorded in the fruit sample of \textit{T.bellerica}. The highest value of 1117.08± 73.69 nutritive value was recorded in the fruit sample of \textit{S.anacardium} and lowest value of 561.90±5.38 was recorded in the fruit sample of \textit{B.frondosa} (Table 2 and Fig 3, 4).

![Figure 3](image-url) - Variation of proximate (%) in different medicinal plants

![Figure 4](image-url) - Variation of Nutritive value (Cal/100gm) in different medicinal plants
Table 3: Comparison of Recommended Dietary Allowances (RDA) with proximate, nutritive value and elemental composition of medicinal plants of Shimoga district

<table>
<thead>
<tr>
<th>Sl No</th>
<th>Factors</th>
<th>T.chabula</th>
<th>T. bellerica</th>
<th>E. officinalis</th>
<th>S. trifolius</th>
<th>S. anacardium</th>
<th>B. frandonia</th>
<th>RECOMMENDED DIETARY ALLOWANCES</th>
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<td>V</td>
<td>Nutritive value (Cal/100mg)</td>
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<td>958.34</td>
<td>882.90</td>
<td>1174.08</td>
<td>1117.08</td>
<td>561.90</td>
<td>55 to 60%</td>
<td>55 to 60%</td>
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</table>
Further, Obajinwa et al. (2002) said that mineral contents of plants varies according to composition of soils in which plants is grown. When the heavy metals are not recorded in the samples suggesting that the herbs are collected from unpolluted natural habits and reflect the natural levels of heavy metals. In the present study, the heavy metals were not recorded in the plant samples except Terminalia chebula and Terminalia bellerica in which trace of lead and cadmium was recorded. The similar observations were also made by Sarala (2006) and Shashikumar (2006) in the plants of Withania somnifera and different species of Dioscorea which were collected from the different habitats of different climatic conditions.

Nutritive values of the medicinal plants which were studied ranged between 294 cal/100 gm and 331 cal/100 gm in the fruit samples of Emblica officinalis and Terminalia chebula respectively. Donatus Ebere (2005) studied the components of the nutrients, carbohydrates crude fibres, moisture content and crude protein of ripen fruits of Dennettia tripetala and reported that the calorvic value of 480.24 gm, cal/100 gm of fresh fruits. Simultaneously, mineral content of calcium (1.8%), phosphorus (0.33%), potassium (2.50%) and magnesium (0.42%) and they have justified uses of Dennettia tripetala fruits as food and a drug in herbal medicine in south eastern Nigeria. Further, Indrayana et al. (2005) tried to evaluate nutritive values of seeds of Nelumbo nucifera, Embelia ribes, Eugenia jambolana and Artocarpus heterophyllus and they have classified plants into food and fodder uses. Deepak Dhyani (2007) attributed studied basic nutrients of Hippophae rhamnoides population from Uttara Khand, India and said that the nutritive fruit values of fruits and seeds of Seed buckthorn varied in different locations and also emphasized the importance of nutritional values for the development of organic foods and nutritional industries in the state of Uttara Khand. Therefore, the present investigation, Terminalia chebula, Terminalia bellerica and Emblica officinalis are used in the proportion of Triphala an ayurvedic formulation (Naik et al., 2006). Hence, the present study determination of elemental composition and nutritive values of Terminalia chebula, Terminalia bellerica and Emblica officinalis have important connotation in light of upcoming ayurvedic medicines and plant based pharmaceutical industries of India.

CONCLUSION

The six plants which are studied in the present investigation, the fruits of Terminalia chebula, Terminalia bellerica and Emblica officinalis are used in the preparation of triphala, an ayurvedic formulation. The plants have rich in the sources of proximate, nutritive value and elements composition. The results were compared with RDA values all the components are rich in the fruit samples. The fruits of Semicarpus anacardium and Sapindus trifoliatus which belongs to same family Anacardaceae are used in the preparation of ayurvedic formulations and the study is important connotation in light of upcoming ayurvedic medicines and plant based pharmaceutical industries of India.

Acknowledgements

The authors thank the Chairman, Department of Applied Botany, Kuvempu University, Shankaraghatta, Shimoga.

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