



RESEARCH ARTICLE

A STUDY ON SELECTED INDIVIDUAL TREE CANOPY OF *BAUHINIA PURPUREA*,  
LINN;- IN URBAN GREENING

\*Arul Sheeba Rani, M. and Mary Josephine, R.

Department of Botany, Nirmala College for Women, Coimbatore, India

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ABSTRACT

Urban greening refers to any vegetation effort including the planting of trees, shrubs, grass or agricultural plots whose design is intended to improve the environmental quality, economics opportunity or aesthetic value associated with a cities landscape. For the present study tree *Bauhinia purpurea*, linn.; were selected for the physico-chemical parameters of tree canopy soil, mineral profile of the litter formed by the selected tree canopy, microbial flora of the selected tree canopy soils were analyzed. Hence, the present study the aim is to improve our quality of life in an increasingly densely populated, fast-living world. People have to find then way back to natural and green open spaces that become more and more important for our personal development, wellbeing and recreation due to increasing urbanization.

INTRODUCTION

Impervious cover plays an important role in the landscape, particularly in urban areas. These surfaces such as roads, buildings, sidewalks and parking lots facilitate transportation and provide shelter. Trees, forests, open spaces, rivers and streams and associated natural resources improve our quality of life and provide us with a sense of community, improve our individual and community self-esteem and promote our physical and mental well-being. The enhancement of urban green spaces or urban green forests is one of the ways, which has the potential to mitigate the adverse effects of urbanization economic or environmental costs. Urban greening is an integrated approach to the planting, care and management of all vegetation in cities, towns, townships and informal settlements in urban areas. Urban green spaces play a significant role for people to have social contacts or find rest in order to achieve this inner harmony and well being.

MATERIALS AND METHODS

Tamil Nadu is one of the 28 States of India. Its capital is Chennai (formerly known as Madras) the largest city. Nirmala college academic campus is located in the southern parts of the Western Ghats.

\*Corresponding author: Arul Sheeba Rani, M.

Department of Botany, Nirmala College for Women, Coimbatore, India.

The temperature during both summer and winter varies between 28° c to 34° c. Soil in this area is red loamy soil which is more fertile than sandy soil. Its porosity allows high moisture retention and air circulation.

Collection of selected tree sample

For the present study *Bauhinia purpurea*, linn.; were selected in the Nirmala college campus to find out the morphology and propagation of the selected tree, physico-chemical parameters of the tree canopy soil, mineral profile of the litter formed by the selected tree canopy, microbial flora of the selected tree canopy soils were analyzed.

Taxonomic Position

Division : Phanerogams  
Class : Dicotyledons  
Subclass : Polypetalae  
Series : Calyciflorae  
Order : Rosales  
Family : Fabaceae  
Subfamily : Caesalpinieae  
Genus : *Bauhinia*  
Species : *B. purpurea*, Linn.;

**Bauhinia purpurea**Linn: is a native of South eastern Asia from India to China and is planted in Florida.It is commonly called Hong Kong Orchid Tree. It is a small to medium-sized

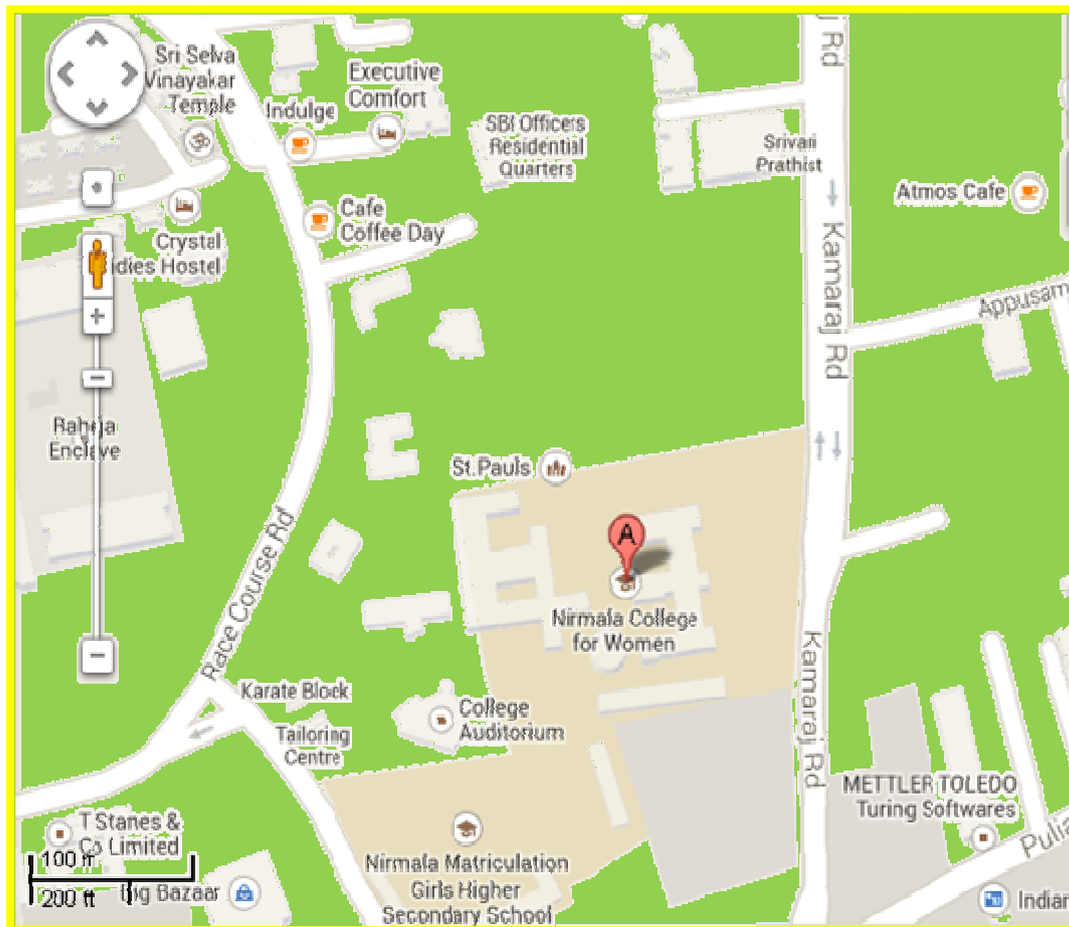


Plate 1. Location Map

deciduous fast-growing shrub or tree with a round, symmetrical, moderate dense crown to 10 m tall. It is used in traditional medicine as treatments for many ailments, such as ulcers, wounds, even cancer-fighting effects, swollen glands, and stomach tumours. It is extensively applied in glandular diseases and poulticing to reduce swelling and bruises and to ripen ulcerations also boils as a poison antidote and as cough medicine. Decoctions of various plant parts are taken internally as an astringent, febrifugal, antidiarrhoeal, anti-inflammatory and antidyenteric remedy.

#### A. Morphological characteristics of the selected tree and propagation

Morphological characters of the selected tree species were recorded. The selected trees total height and width. Leaf, leaflet, flower, fruits - size and colours were measured.

#### Biodiversity of the selected tree

Biodiversity of species such as Ants, Crow, Sparrow, Pigeon, Dragon fly, Mynah, Butterflies, Lac insect, Lizards, Calottes, Chameleon, Spider, Worms, Honey comb, Honey bee, Wasp, Parrots, Grasshopper, Sparrow were observed and recorded during the study period.

#### Average annual litter of dried leaves and logs of the selected tree canopy

The litter of dried leaves and logs of the selected tree canopy were collected throughout the year and the average annual fallings were calculated.

#### Microbial analysis

##### Collection of the selected tree canopy soil sample

The tree canopy soil samples were collected during the year, 2014-2015. Soil with litter formation and ground vegetation from the selected tree canopy of *Albizia* were collected separately in sterile bags, air dried and sieved for further analysis. Barren land soil, taken from the same campus was kept as control. Soil was taken from the depth of (0-15 cm depth). Soil samples were packed in sterile bags and used for further analysis.

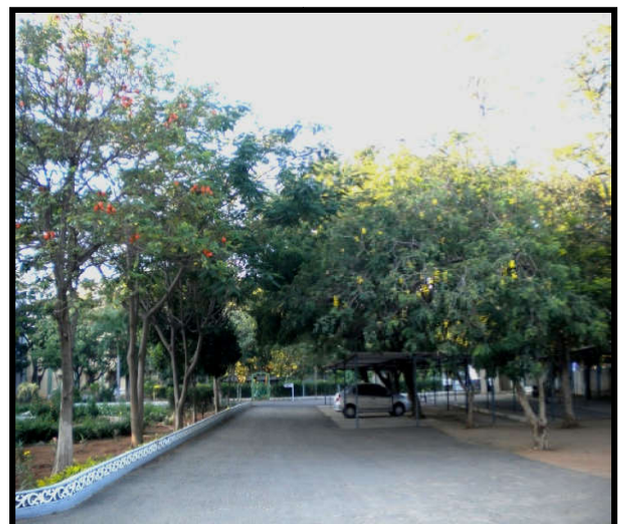


Plate 2. Study Area



Plate 3. Habit

### Isolation and culture of microorganisms

#### Preparation of nutrient medium: Potato-Dextrose Agar (PDA)

120 gms of freshly peeled potato is taken in to a flask and 150 ml of water is added to it. It is boiled for 10 minutes. Then the potato extract is taken and its volume is made up to 150 ml by adding distilled water. To this extract, 7.5 gms of Dextrose is added and thoroughly mixed. Then the solutions were poured in a 500 ml flask and stirred thoroughly. This content is heated in a water bath to dissolve the agar. This medium is dispensed in culture petridishes and kept in laminar air flow for solidification.

#### Serial dilution method

For the enumeration of microbial population a set of ten selected soil samples (0-15 cm depth) were collected. Soil microbial communities have relied on culturing techniques using PDA (Potato Dextrose Agar) medium. Serially diluted samples were inoculated on petridishes containing PDA medium and incubated in the laboratory for 5 days at 30°C (Kanika Sharma, 2007).

#### Identification of Bacteria

An average volume of bacterial cell is 1 cubic micron. They are smallest forms among bacteria. After division the cells may either separate from each other or may remain joined together to form groups of two cells in *Diplococcus*, a tetrad of four cells in *Micrococcus tetragenus* and a chain of cells in *Streptococcus* (Bergey, 1957).

#### Identification of Fungus

The smear was simple stained to study the morphology of the cells. Basic stain for simple staining Safranin is used for identifying microbes and the data's were recorded. For each experiment replicas were repeated (Mani *et al.*, 2004).

### Physicochemical parameters

Physicochemical parameters of the select tree canopy, litter and barren soils were analyzed.

#### pH of the soil

Part of the moist soil samples were air dried and sieved to obtain fine soil samples (2 mm). The pH of the medium, if found to be acidic, is brought to the required pH by adding 0.1 (N) NaOH drop wise and testing with pH paper after thoroughly mixing with a glass rod.

#### Moisture content of the soil

Moisture content of the selected tree canopy litter samples were calculated and expressed in percentage (Conventional oven method ASTM, 2001).

#### Water holding capacity and temperature of the soil

Water holding capacity and temperature of the soil were analyzed as per the standard method.

#### Mineral profile of the selected tree canopy soil samples

Mineral like Potassium, Phosphorus, Calcium, Magnesium, Iron and Sodium were analyzed in the standard laboratory by employing Atomic Absorption Spectrophotometer by following the method of Issac and Johnson (1975) and the results were recorded.

#### Estimation of calcium and magnesium (Jackson, 1967)

5ml of triple acid digested extract was taken in a China dish. To this 10 ml of 10% NaOH and 0.1g of Murexide indicator powder were added and titrated against 0.02 N versenate (19 g of EDTA was dissolved in 5liters of distilled water) and standardized against 0.2 N Na<sub>2</sub> CO<sub>3</sub> solution and adjusted until the colour changes from red to violet.

#### Calcium and Magnesium

5ml of triple acid digested extract was taken in a China dish, to this 10 ml of ammonium chloride - ammonium hydroxide buffer pH 10 and few drops of Eriochrome Black T indicator were added and titrated against 0.02N versenate solution until the colour changes from red to blue.

#### Estimation of Sodium and Potassium

Sodium and potassium were estimated by using Flame Photometer, Model-EFL. The sodium and potassium contents were calculated by referring to the calibration curves of sodium and potassium, respectively, and expressed as mg/100 g on dry weight basis.

#### Phosphorus estimation (Dickman and Bray, 1940)

One ml of triple acid digested extract was pipetted into 100 ml volumetric flasks. To this 50 ml glass distilled water was added, followed by 5 ml of ammonium molybdate sulphuric acid reagent Solution A was added slowly with constant stirring to solution B and the volume was made up to 100 ml with glass distilled water). Blue colour was developed by adding six drops of 2.5% stannous chloride solution.

**Table 1. Comparative Morphological characters, Propagation and the Biodiversity of the selected tree sample**

Sample	Tree	Height in (m)	Breadth in (m)	Leaf		Inflorescence	Flower colour	Fruit		Seed shape and colour	Propagation	Biodiversity
				Type	Shape							
<i>Bauhinia purpurea</i>	Deciduous, Evergreen and Shrubs	07.05	01.65	Smooth and elliptic,	Rounded shallow to Cordate	Raceme in terminal panicles	Rich purple, Dark Pink	Rounded, flat	Elongated dehiscent pods	Shiny-brown,	Seeds	Ants, Butterflies

**Table 2. Morphology of the Leaf/ Leaflet length**

Sample	Simple/compound	Leaf length in (cm)	Leaflet length in (cm)	Leaf/ Leaflet of the selected trees
<i>Bauhinia purpurea</i>	Simple smooth and elliptic,	11.00	0.00	

**Table 3. Morphology of the inflorescence and flower of the selected tree**

<i>Bauhinia purpurea</i>	Raceme in terminal panicles	Rich purple, Dark Pink	08.09	
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**Table 4. Dehiscent and indehiscent seeds of the selected tree**

Sample	Pod
<i>Bauhinia purpurea</i>	Dehiscent/ Indehiscent Dehiscent

**Table 6. Average annual litter of dried leaves and logs of the selected tree canopy**

Sample	January- March (gm)	April - June (gm)	July- September (gm)	October-December (gm)	Average annual litter of the selected tree canopy in (%)
<i>Bauhinia purpurea</i>	177.33	417.87	260.00	453.00	3.27

**Table 7. Enumeration of the Bacterial colony of the selected tree canopy soil**

Sample	Number of Bacterial Colony								
	Day 1			Day 2			Day 3		
	$10^{-3}$	$10^{-6}$	$10^{-9}$	$10^{-3}$	$10^{-6}$	$10^{-9}$	$10^{-3}$	$10^{-6}$	$10^{-9}$
Control	3	3	2	5	4	3	5	7	6
<i>Bauhinia purpurea</i>	6	4	3	7	5	4	9	7	6

**Table 8. Bacteria present in the selected tree canopy soil**

Sample	Bacteria		
	$10^{-3}$	$10^{-6}$	$10^{-9}$
Control	<i>Streptococcus sps</i>	<i>Staphylococcus sps</i>	<i>Streptococcus sps</i>
<i>Bauhinia purpurea</i>	<i>Pseudomonas sps</i>	<i>Staphylococcus sps</i>	<i>Stepotmycetessps</i>

**Table 9. Enumeration of Fungal colony of the selected tree canopy soil**

Sample	Number of Fungal Colony								
	Day 1			Day 2			Day 3		
	$10^{-3}$	$10^{-6}$	$10^{-9}$	$10^{-3}$	$10^{-6}$	$10^{-9}$	$10^{-3}$	$10^{-6}$	$10^{-9}$
Control	-	-	-	3	3	2	3	3	2
<i>Bauhinia purpurea</i>	-	-	-	1	1	2	3	2	2

**Table 10. Fungus present in the selected tree canopy soil**

Sample	Fungi		
	$10^{-3}$	$10^{-6}$	$10^{-9}$
Control	<i>Aspergillus niger</i>	<i>Aspergillusglaucus</i>	<i>Aspergillus niger</i>
<i>Bauhinia purpurea</i>	<i>Aspergillusglaucus</i>	<i>Aspergillusglaucus</i>	<i>Aspergillusglaucus</i>

**Distribution of Microbes present in the selected individual tree canopy soil (plate -4)****Table 11. Moisture content and pH of the selected tree canopy soil**

Sample	Fresh weight (gm)	Dry weight (gm)	Moisture content(%)	pH
Control	20	18.86	5.7	5.7
<i>Bauhinia purpurea</i>	20	18.57	7.2	5.8

**Table 12. Mineral profile of the selected tree canopy soil**

Sample	Potassium (%)	Phosphorus (%)	Calcium (%)	Magnesium (%)	Iron (%)	Sodium (%)
Control	0.39	0.10	0.31	0.081	0.048	0.18
<i>Bauhinia purpurea</i>	0.26	0.10	0.52	0.11	0.009	0.49

**Table 13. Moisture content and pH of the selected tree canopy litter**

Sample	Fresh weight (gm)	Dry weight (gm)	Moisture content(%)	pH
<i>Bauhinia purpurea</i>	177.33	100.00	43.60	6.2

**Table 14. Mineral profiles of the selected tree canopy litter**

Sample	Potassium (%)	Phosphorus(%)	Calcium(%)	Magnesium(%)	Iron(%)	Sodium(%)
<i>Bauhinia purpurea</i>	855	228	990	260	25	42

The total volume was made up to 100 ml. The intensity of the blue colour was measured at 650 nm in a spectrophotometer. The phosphorus content present in the sample was calculated by referring to a standard curve of phosphorus and expressed as mg/100 g on dry weight basis.

#### **Estimation of iron by atomic absorption spectrophotometer (Issac and Johnson, 1975)**

By feeding the sample to an Atomic Absorption Spectrophotometer the iron content was estimated at 246.8 nm wavelength and the readings were expressed in mg/100g of sample on dry weight basis.

#### **V. Analysis of the selected tree canopy litter formed by the selected samples**

##### **Collection of tree canopy litter samples**

From a composite of litter fall, the fallen fresh/dried leaves, wood logs, flowers, fruits and seeds were collected under the canopy of the ten trees separately and shade dried, packed in sterile bags then powdered and lumped in a composite of sample for chemical analysis. The maximum litter fall of various seasons during the year 2014 (January-March, April-June, July-September, October-December) were analyzed.

##### **pH and moisture content**

pH and moisture content of the litter were analyzed as per the standard methods.

##### **Mineral analysis of the selected tree canopy litter samples**

Mineral profiles of the litter formed by the selected tree canopy, the fallen fresh/dried leaves, wood logs, flowers, fruits and seeds were powdered and kept in airtight container then the mineral profiles were analyzed and the mineral profile of the selected tree canopy soil and litter samples were experimented and recorded by following standard methods of (Association of Official Agricultural Chemists) AOAC, (1990).

## **RESULTS AND DISCUSSION**

Comparative morphology of the selected trees, leaves, inflorescence, flower, fruit, pod (dehiscent/indehiscent) and its propagation, Micro and Macrobial biodiversity were observed and represented in the following Tables.

## **Conclusion**

India is urbanizing at a very fast pace. The enhancement of urban green spaces or urban green forests is one of the ways, which has the potential to mitigate the adverse effects of urbanization economic or environmental costs. The research on urban greening is very meagre particularly in India. Planting tree is the need of the hour. As tree grows their component value increases. Healthy trees contribute to the overall value of its properties to the society. Urban green areas contribute to maintaining and expanding the biological base for diversity that is essential to human survival in to the millennium. Hence, the study on selected individual tree canopy of the soil and litter in urban greening to enrich the urban soil and to promote plant growth to the urban environment

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