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# Histochemical investigation of different organce of genus sesbania of marathwada region in maharashtra

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#### Abstract

The histochemical studies of leaves and wood of Sesbania grandiflora, Sesbania bispinosa and Sesbania cannabina are medicinally important plants of Marathwada region in Maharashtra. For histochemical studies the free hand sections of leaves and wood were taken and treated with the respective reagent in localize components, *viz.* starch, protein, tannin, saponin, fat, glucosides and alkaloids in the tissues.

Keywords: Histochemistry, starch, protein, tannin, saponin, fat, glucosides and alkaloids

# INTRODUCTION

Histochemistry is the branch of histology dealing with the identification of chemical components of cells and tissues. Starch deposition occurs widely in the plant body, but the particularly common places of its accumulation are seeds, the parenchyma of the secondary vascular tissues in the stem and root, tubers, rhizomes and corn(Kadam , 1999). Starch and proteins are the principal ergastic substances of the protoplast (Kuster, 1956). Tannin is the heterogeneous group of phenol derivatives, usually related to glucosides (Vaidya, 1972). Tannins are particularly abundant in the leaves (xylem) of many plants (Kadam et al., 1996). Saponin are the rare occurrence. Fats are widely distributed in the plant body and they probably occurs in small amount in every plant cell (Seifriz, 1934).Fats are common reserve material in seeds, spores and embryos in meristematic cells. Glucosides are the degradation product of the carbohydrates. Alkaloides are the degradation product of protein.

Many woody plants contain medicinally important secondary product (Dhar *et al.*,1968). Therefore, we have attempted to histochemical investigations of different plant parts of *Sesbania grandiflora*, Sesbania bispinosa and *Sesbania cannabina*, three medicinal plants of Marathwada region in Maharashtra. Free hand sections were taken for the histochemical studies. Sections are treated with the respective reagent to localize components, viz. starch, proteins, tannin, saponin, fat glucosides and alkaloids in the tissues (Johansen, 1940).

## MATERIALS AND METHODS

Temporary and permanent mounts of sections were employed

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Tel : +91-9423743987 Fax:+91-253-2577341 Email: drvbkadam@yahoo.com for the test of histochemical studies. For study of isolated different tissues, small pieces of material were macerated in Jeffery's fluid (Johansen, 1940). For the histochemical studies free hand sections of the organs to be studies, were taken and treated the respective reagent to localize component, Viz. starch, protein, tannin, saponin, fat, glucosides and alkaloids in the tissues (Johansen, 1940).

### Starch

0.3 g of iodine and 1.5 g of potassium iodide were dissolved in 100 ml of distilled water. A drop of the solution was added on the section, washed water and observed under microscope.

#### Protein

Saturated aqueous solution of picric acid is an excellent precipitating agent for protein, staining them an intense yellow. It was allowed to react with the reagent for 24 hours. b)Dilute eosin, stains protein red. c) To localize protein, reagent was prepared by mixing 0.1 g potassium ferro cyanide dissolved in 20 ml water and 100 ml glacial acid. Section was kept in for an hour. They section were washed with 60% alcohol and few drop of aqueous Fecl<sub>3</sub> were added .Blue color indicates the presence of proteins.

#### Tannin

Sections were treated with dilute acidic Fecl<sub>3</sub> solution (0.5% to 1 % of ferric chloride in 0.1 N HCL); mounted in clove oil and observed under microscope for the presence of tannins. 10% aqueous Fecl<sub>3</sub> plus little Na<sub>2</sub>co<sub>3</sub>; blue green colour is given by tannin.

## Saponins

Sections were placed directly in one drop of concentration  $H_2So_4$  on a slide, which gives a characteristic sequence of colour reactions, beginning immediately with yellow, changing to red within 30 minutes and finally becoming violet or blue green in a short time. To determine localization of the saponin, sections were put in saturation barium hydroxide solution for about 24 hours. Sections were washed with calcium chloride, the placed in potassium

dichromate. Yellow colour indicated the presence of saponins.

# Fat

0.5 g of dye, Sudan III or Sudan IV was dissolved in 100ml of 70% alcohol. Sections were kept in the stain for 20 minutes, rinsed quickly with 50% alcohol and mounted in glycerin for observations. Blue, red, pink, precipitate indicated the presence of fat.

# Glucoside (Goignard's test)

Section were immersed in 1% of aqueous picric acid for 30 minutes, washed with water and placed in a drop of 10% aqueous sodium carbonate. A red colour of the section with hydrochloric acid reveals the of Glucosides. For the localization, section were placed in solution composed of 20 parts of 20% aqueous KOH and 80 parts of 90% alcohol for few minutes. In a small watch glass, mixture of 2.5% aqueous Feso<sub>4</sub> and 20% aqueous Fecl3 solution taken in equal proportion was heated to boiling and then the sections were transferred to a slide holding a drop of 20% hydrochloric acid. A deep blue precipitates indicates indicated the presence of glucosides.

## **Test for Alkaloids**

Transverse sections of the different plants were treated with the following with the following alkaloid reagent.

a) Mayer's Reagent

Potassium mercuric iodide solution; 13.55g of  $HgCl_2$  and 50 g of KI, were dissolved in one liter of distilled water. Presence of grey colour in the section reveals the presence of alkaloids.

b) Wagner's Reagent

1gm iodine and 2g potassium iodide were dissolving in 50ml of distilled water. Presence of golden yellow colour reveals the presence of alkaloids.

## **RESULTS AND DISCUSSION**

Histochemical localization in different organs of the taxa under study was made, using methods described elsewhere. The initial presentation gives details about the occurrence of erastic content or secondary metabolites, *viz.*, starch, protein, fat, tannin, saponin, glucoside and alkaloids in leaves and Wood.

**Starch :** Starch is the principal ergastic substance of the protoplast. Starch is composed of long chain molecules, whose basic units are anhydrous glucose residues of the formula  $C_6H_{12}O_5$ . Starch has an ordinary arrangement of molecule and, therefore, shows optical anisotropy and double refraction. In starch granules the molecule is radically arranged, therefore, in polarized light a cross pattern is seen. The morph metric Variation of starch grain is so extensive that they may be used taxonomically and pharmacognostically up to a limited extent (Kuster, 1956).

Starch deposition occurs widely n the plant body, but the particularly common places of its accumulation are seeds, the parenchyma of the secondary vascular tissue in wood and roots, tuber, rhizome and corms. In the present work, for the taxa under study, starch was present in leaves and wood of all the taxa, *viz.*, *Sesbania grandiflora* (Table 1) (Sesbania bispinosa (Table.2), *Sesbania cannabina* (Table 3)

Sr. No.	Ergastic content	Reaction		Localization		
		Leaves	Wood	Leaves	Wood	
1	Starch	*ve	*ve	Scattered cells of mesophyll, Mid -rib pith parenchyma,	Cortical parenchyma, Medullary rays, Vascular bundle, and Pith parenchyma	
2	Protein	-do-	-do-	Epidermis, Scattered cells of mesophyll,	Epidermis, Scattered cells of Cortex.	
3	Tannin	-ve	-do-	Scattered cell of mesophyll	Scattered cells of Cortex and Pith parenchyma	
4	Saponin	+ve	-do-	Scattered cell of mesophyll	Epidermis, Scattered cells of Cortex parenchyma, and Pith	
5	Fat	-do-	-do-	Upper and lower epidermis, Scattered cells of Mesophyll cells and Mid –rib	Cortical parenchyma, Medullary rays, Scattered cells of Pith parenchyma.	
6	Glucoside	+ve	*Ve	Lower epidermis, Scattered cell of mesophyll	Epidermis and Scattered cells of Cortex and Pith parenchyma	
7	Alkaloids	-				
	a) Mayer's reagent	*ve	*ve	Epidermis and cortical cells	Cortex, Xylem parenchyma, and Pith parenchyma.	
	b)Wagner's reagent	-do-	-do-	Upper and lower Epidermis, Mid – rib parenchyma.	Epidermis, Cortical parenchyma, Medullary rays and Vascular bundle and Pith parenchyma	

#### Table 1-Histochemical test for fresh section of leaves and wood of Sesbania grandiflora

**Protein:** Protein are the major constituents of the living protoplast, but they also occur as temporarily inactive erastic substance. Erastic protein is knows as a storage material and is found deposited in amorphous and or crystalline forms. Like starch

and cellulose, crystalline protein combine crystalline and colloidal properties, therefore, the individual units of this material are spoken of as crystalloids (meaning crystal like) rather than as crystals.

This is also present in all the taxa under investigation. Protein

were observed in the upper and; lower epidermis, scattered cells of mesophyll of leaves, and cortical parenchyma in the wood of

Sesbania grandiflora (Table. 1) (Sesbania bispinosa (Table. 2), Sesbania cannabina Linn (Table .3)

Table 2. Histochemical test for fresh section of leaves and wood of Sesbania bispinosa

Sr. No.	Ergastic content	Reaction		Localization	
		Leaves	Wood	Leaves	Wood
1	Starch	*Ve	*ve	Upper and lower epidermis, Scattered cells of Mesophyll, Mid –rib Parenchyma, Pith parenchyma	Xylem and Phloem parenchyma and Scattered cells of Medullary ray. and Scattered cells of Cortex
2	Protein	-do-	-do-	Upper and lower epidemis, cells of mesophyll cells, Xylem parenchyma, Scattered cells of medullary rays and Pith parenchyma.	Epidermis, Scattered cells of cortex parenchyma, Xylem and Phloem,Scatteredcells of medullary rays, Pith parenchyma
3	Tannin	-do-	-do-	Scattered cells of mesophyll and Pith cells	Scattered cells of medullary ray, and Pith parenchyma
4	Saponin	-do-	-do-	Mesophyll cells , Pith parenchyma,	Scattered cells of cortex parenchyma and Pith region, Xylem parenchyma
5	Fat	-do-	-do-	Scattered cells of epidermis , Mesophyll cells and mib- rib Pith parenchyma	Vascular bundle and Scattered cells of medullary rays and Pith parenchyma
6	Glucoside	*Ve	*Ve	Epidermis and cortical cells	Epidermis, Scattered cells of cortex parenchyma
7	Alkaloids	-do-	-do-		
	a)Mayer's reagent	*ve	*ve	Upper and lower epidermis, Scattered cells of mesophyll cells.	Hypodermis, Xylem parenchyma ,Pith
	b)Wagner's reagent	-do-	-do-	Upper and lower epidermis , Scattered cells of mesophyll cells, Medullary rays , Mid-rib, Pith parenchyma	Epidermis, Scattered cells of cortical parenchyma, Medullary rays, and Vascular bundle

Table 3. Histochemical test for fresh section of leaves and wood of Sesbania cannabina

Sr.	Ergastic	Reaction		Localization	
No.	content	Leaves	Wood	Leaves	Wood
1	Starch	⁺ve	*ve	Upper and lower epidermis , Mesophyll cell, Cortical cells, Pith parenchyma	Medullary rays, Cortical parenchyma
2	Protein	-do-	-do-	Scattered cells of cortex, Mesophyll cells and Pith	Epidermis ,Cortical parenchyma, Pith parenchyma
3	Tannin	-do-	-do-	Scattered cells of mesophyll, Mid-rib pith parenchyma.	Vascular bundle and Scattered cells of medullary ray.
4	Saponin	-do-	-do-	Upper and lower epidermis and Mid-rib pith	Scattered cells of cortex and Medullary rays, and Pith parenchyma
5	Fat	- do-	-do-	Mesophyll ,cortical and in pith	Scattered cells of pith ,Cortex, and Medullary rays
6	Glucoside	-do-	-do-	Epidermis and cortical cells	Scattered cells of cortex , Medullary rays and Vascular bundle
7	Alkaloids				
	a)Mayer's reagent	*ve	*ve	Upper epidermis Scattered cells of mesophyll, Mid rib pith parenchyma	Scattered cells of cortex, and Vascular bundle

**Tannin:** Tannin is a heterogeneous group of phenol derivatives, usually related to glucosides. Tannins are particularly abundant in the leaves of much plant; in the xylem, in the testa of seeds and in pathological growth like galls (Kuster, 1956). No tissue, however, appears to lack tannins entirely. Sometimes tannins containing cells are conspicuously associated with a vascular tissue terminates beneath storage tissue or secretary cells of nectarines. The monocotyledons are notably poor in tannins.

Tannins also show distributions, occurring mostly in epidermis, mesophyll cortical as well as parenchymatous tissue, associated with conductive tissue. Tannins were observed in the leaves of *Sesbania grandiflora* (Table1) *Sesbania bisponisa* (Table 2), *Sesbania cannabina* Linn (Table 3)

Saponin: The saponin is of rare occurrence and wherever present, they apparently remain to one or two organs. Saponin were observed in the mid-rib parenchyma of leaves and cortex and pith parenchyma of wood Sesbania grandiflora (Table 1) (Sesbania bispinosa (Table 2), Sesbania cannabina Linn (Table 3). Saponin were observed in the cells of mesophyll and xylem parenchyma of wood of Sesbania grandiflora (Table1) (Sesbania bispinosa (Table 2), Sesbania cannabina Linn (Table 3)

**Fat:** Fat are widely distributed in the plant body, and they probably occur in small amounts in every plant cell. The term fat may be used to described not only the fats proper (that is, ester of fatty acids with glycerol), but also related substances grouped under the name of lipids.

As protoplast inclusion, fats are common reserve material in seeds, spores and embryos in meri woodatic cells and occasionally in differentiated tissue of the vegetable body. They occur as solid bodies or, more frequently, as fluid droplets of various size either dispersed in the cytoplasm or aggregated in large masses fatty substance are thought to be elaborated directly by the cytoplasm and also by leucoplast. In taxa under study, fat was found in cells of mesophyll and phloem parenchyma (leaves and wood) of Sesbania

grandiflora (Table1) (Sesbania bispinosa (Table 2), Sesbania cannabina Linn. (Table 3)

**Glucoside**: Glucosides are the degradation production of carbohydrates glucosides were observed in the epidermis ,pith parenchyma of leaves vascular bundles and scattered cells of medullar ray of wood Sesbania grandiflora (Table1) Sesbania bispinosa (Table 2), Sesbania cannabina Linn (Table 3)

**Alkaloids**: Alkaloids are degradation of protein they were investigated by using two methods, namely; Mayer's reagent and Wagner's reagent. In Mayer's reagent alkaloids were observed in the scattered cells of mesophyll of leaves and pith parenchyma of wood .In Wagner's reagent, alkaloids were found in the cells of mesophyll and cells of cortex parenchyma and pith parenchyma of wood of Sesbania grandiflora (Table 1) (Sesbania bispinosa (Table 2), Sesbania cannabina (Table 3)

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