EVALUATION OF PHARMACOGNOSTICAL AND PHYTOCHEMICAL PROPERTIES OF THE LEAVES OF Psidium guajava Linn - NAGPUR SEEDLESS VARIETY

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INTRODUCTION

Psidium guajava Linn commonly called as poor man apple. The leaves of P. guajava really do not have any match as a cheap natural and easily available plant. It is traditionally known to be useful for the treatment of wide panel of diseases like ulcers, wounds, astrignent, antiemetic, cholera, epilepsy etc[1]. Leaf is traditionally used for antispasmodic, anodyne, febrifuge[2], scurvy[3], malaria[4], antibiotic[5], antifungal[6-8], antifungal, anti-inflammatory[9-14], antidiabetic[10,11], anti-hypertensive[12], analgesic[13], hepatoprotective[14] and anti-coagulant[21].

It was reported that fresh leaves contains: Guajavarin, isoorcetrin, hyperin, quecetrin, quercetin 3-o gentiobioside[22]. Leaves also contains two triterpenoids, guavaonic acid and guava coumaric acid along with six known compounds 2 alpha hydroxy ursolic acid, jacoumaric acid, isonomucomaric acid, asiatric acid, ileopal D and beta-sitosterol – 3-o – beta D glucopyranoside[23]. In short, there is good level of traditional and experimental evidences to support various claims and advantages of this widely available plant. An investigation to explore its pharmacognostic examination is inevitable. Hence, in this work we report an attempt on microscopic evaluation, physico-chemical determination and phytochemical screening for the standardization and quality assurance purposes of this cultivar.

MATERIALS AND METHODS

Chemicals

Formalin, acetic acid, ethyl alcohol, chloral hydrate, toludine blue, phloroglucinol, glycerin, hydrochloric acid and all other chemicals used in this study were of analytical grade.

RESULTS

Plant collection and authentication

The leaves of the healthy plant Psidium guajava Linn. (Nagpur seedless) selected for our study was collected from Horticulture Department, Madurai, Tamil Nadu, India and was authenticated by Dr.Stephen, Department of Botany, American college, Madurai and Dr. P. Jayaraman, Director of Plant Anatomy Research Institute, Tambaram, Chennai, Tamil Nadu, India.

Macroscopic analysis

Macroscopic observation of the plant was done. The shape, size, surface characters, texture, colour, odor, taste etc was noted[24].

Microscopic analysis

Transverse section midrib region of fresh leaf pieces were cut and fixed in FAA and then dehydrated by employing graded series of ethyl alcohol and tertiary butyl alcohol[25]. Sections were taken using microtome. Permanent mount was prepared using saffranin fast green double staining technique[26]. In order to supplement the descriptive part the photomicrographs in different magnifications of all necessary cells and tissues were taken with NIKON Coolpix 8400 digital camera and Labphot 2 microscopic unit.

Powder microscopy

Coarse powder of the leaf was used to study the microscopical characters of the leaf powder[27,28].

Physicochemical analysis

Total ash, acid insoluble ash, water soluble ash, loss on drying, extractive values and leaf constants such as vein islet numbers, vein terminal number, stomatal number and stomatal index, palisade ratio were determined[29-31].
Preliminary phytochemical screening

Preliminary phytochemical screening was carried out to find out the presence of various phytoconstituents using standard procedure [32-33].

RESULTS

Macroscopy

Psidium guajava is a large dicotyledonous shrub or small evergreen tree, generally 3-10m high with many branches and crooked stems (Fig 1). Leaves (5-10cm × 4-5cm) are opposite, simple, stipules absent, oblong – elliptic, dull grey to yellow green with entire margin, obtuse to bluntly acuminate apex and rounded to subcuneate base with short petiole (Fig 2). Flowers are white, borne singly or in small clusters, 2-3 cm wide, with 4 or 5 white petals which are quickly shed, and a prominent tuft of perhaps 250 white stamens. Fruit is small, 3 to 6 cm long, pear-shaped, reddish-yellow when ripe.

Microscopy of the leaf

Transverse section (T.S) of the leaves through the midrib showed the following tissue systems.

Shape: Leaves are dorsiventral with prominent midrib, 1.1mm thick, wide concave adaxial side and horizontally widened with uneven circumference (Fig 3).

Vascular bundle: Vascular strand is broad and bowl shaped, 1.8mm wide, 170µm thick. Xylem elements are thick walled and in long parallel rows. Metaxylem is 25µm in diameter. Phloem is seen in thin dark layer beneath the xylem (Fig 4).
Lateral vein: The lateral vein bundles project prominently below the surface, having thick masses of xylem, small groups of phloem and thick pillar of bundle sheath extensions.

Mesophyll: The palisade cells are two layered with narrow cylindrical compact cells (80µm in height) and short vertically oblong cylindrical spongy mesophyll cells similar to palisade cells.

Ground tissue: Parenchymatous, thick walled circular and compact cells which lacking of tannin.

Epidermis: 210µm thick, even except in the region of the lateral vein. The adaxial epidermis is thin, oblong. The subepidermal layers have wide three rows of polyhedral cells (Fig 5).

Powder microscopy: The analysis of the dried powder of the leaf showed paracytic stomata, multiple layers of wide rectangular cells, epidermal cells with crystals, parenchymal cells without tannin, and fragment of palisade mesophyll, xylem and phloem, fibres.

Physicochemical analysis

Physicochemical parameters were found as follows: total ash 11.05%, water insoluble ash 1.5, water soluble ash 2.77%, ethanol soluble extractive value 18.76%, water soluble extractive value 22.26%, petroleum ether soluble extractive 2.7%, benzene soluble extractive 4.4%, ethyl acetate soluble extractive 5.66%, chloroform soluble extractive 5.24%, loss on drying 9.8%, foreign organic matter was nil. Leaf constants were as follows: vein idlet number 2.3, vein termination number 3.5, stomatal number (lower epidermis) 40, stomatal number (upper epidermis) 32.9, stomatal index (lower epidermis) 19, stomatal index (upper epidermis) 21.

Preliminary phytochemical screening

Preliminary phytochemical screening showed the presence of flavonoids, terpenoids, sterols, tannin, volatile oil, saponins, proteins and amino acids, carbohydrates, reducing sugars, and absence of alkaloids, cyanogenetic glycosides, anthroquinone glycosides, cardiac glycosides, mucilage and fixed oil.

DISCUSSION

Sensory evaluation plays a key role in determining the suitability or denunciation of a crude drug. Organoleptic testing of a crude drug is mainly for qualitative evaluation based on the observation of morphological and sensory profile. In this report, various morphological, microscopic, physicochemical standards have been developed. Hence we have undertaken this study to serve as a tool for developing standards for identification, quality and purity of Psidium guajava leaves.

Adulteration and misidentification of crude drugs can cause serious health problems to consumers and legal problems for the pharmaceutical industries. It can be conducted via a variety of techniques, namely macro and microscopic identification and chemical analysis especially description of microscopic botanical aspects to determine definitively the proper species of plant material while it is still in its non extracted form. The observation of cellular layer morphology or anatomy is a major aid for the authentication of drugs. These characters are especially important for identification of powdered drugs, because in these cases most of the morphological diagnostic features are lost [26]. Microscopic evaluation is one of the simplest and cheapest methods for the correct identification of the source of the materials [24]. The macroscopic and organoepithelial characters of the leaf can serve as diagnostic parameters [30]. Microscopic evaluation showed vascular strand is broad and bowl shaped, 1.8mm wide, 170µm thick. Xylem elements are thick walled and in long parallel rows. The lateral vein bundles project prominently below the surface, having thick masses of xylem, small groups of phloem and thick pillar of bundle sheath extensions. The palisade cells are two layered with narrow cylindrical compact cells (80µm in height) and short vertically oblong cylindrical spongy mesophyll cells similar to palisade cells. The adaxial epidermis is thin, oblong. The subepidermal layers have wide three rows of polyhedral cells.

The ash values are particularly important to find out the presence or absence of foreign inorganic matter such as metallic salts and or silica (earthy matter) [30]. Acid insoluble ash provides information about non-physiological ash produced due to adherence of inorganic dirt, dust to the crude drug. Increased acid insoluble ash indicates adulteration due to dirt, sand (or) soot. The extractive values are primarily useful for the determination of exhausted or adulterated drug and helpful in the detection of adulteration [37]. Phytochemical evaluation and molecular characterization of plants is an important task in medicinal botany and drug discovery [30]. Preliminary phytochemical screening showed the presence of sterols, flavonoids, terpenoids, saponins, volatile oil, protein and aminoacids, reducing sugars, carbohydrates, and absence of alkaloids, fixed oil, mucilage and glycosides. Dried powder of the leaf showed paracytic stomata, three layers of wide rectangular cells, secretory cavity, conical and flagellate trichome, parenchymal cells and fragment of palisade mesophyll.

CONCLUSION

The study of Pharmacognostical features of Psidium guajava Linn. (Nagpur seedless) had shown the standards which will be useful for the detection of its identity and authenticity. The other study viz. physical evaluation, preliminary phytochemical test add to its quality control and quality assurance for proper identification.

Conflict of interest statement:

We declare that we have no conflict of interest.

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