Anatomical Characteristics of African Cherry (Prunus Africana) Medicinal Plant for its Accurate Taxonomic Identification

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Abstract

Background: The genus *Prunus* (Family Rosaceae) comprises over 400 plant species and exhibits vast biodiversity worldwide. Due to its wide distribution, its taxonomic classification is important. Anatomical characters are conserved and stable and thus can be used as an important tool in plant taxonomic characterization. Thus, this study aimed at examining and documenting *P. africana* leaf, stem, and seed anatomy using micrographs and photographs for possible use in identification, quality control, and phylogenetic studies of the species.

Methods: *P. africana* leaves, stems, and seeds were fixed, dehydrated in ascending ethanol series (50–100 %), embedded in Technovit resin, and sectioned using a microtome for mounting histological slides for anatomical observation under a microscope and subsequent description.

Results: The anatomical sections of a young stem revealed a cortex consisting of isodiametric parenchyma cells, druse crystals, primary vascular bundles, and pith. The mature stem bark consisted majorly of rhytidome with periderm densely arranged in multiple layers, a cluster of stone cells, and sclerenchyma. The sections of the leaf were hypostomatic with stomata size ranging between 18.90–(22.34)–26.90 \(\times\) 15.41–(18.40)–21.22 \(\mu m\). The leaf sections showed the presence of characteristic druse crystals, vascular bundles, and mesophyll layers. The pericarp showed the presence of epicarp, mesocarp, and endocarp with a thickness of approximately 350–400, 300–350, and 30–50 \(\mu m\), respectively and a seed testa with a thickness of approximately 50–60 \(\mu m\).

Conclusion: The characteristic morphological and anatomical features observed in *P. africana* leaves, stems, and seeds in this study could provide useful data in taxonomical identification of this species.

1. Introduction

Plant taxonomy plays a critical role in plant diversity assessment, conservation, phytogeographic deductions, optimum utilization, and inferences (Mukherjee, 2014). As such, the events of plant species misidentification could lead to detrimental results. The genus *Prunus* (Family Rosaceae) comprises over 400 plant species and displays vast biodiversity globally, although only approximately 98 species of the genus are of great importance, including *P. domestica* (Linn.), *P. persica* (L.) Batsch, *P. amygdalus* (L.) Batsch, *P. cerasoides* (D.) Don, *P. armeniaca* (Linn.), and *P. africana* (Hook f.) Kalkman (Biswajit et al., 2011). African cherry, also known as Red stinkwood or African almond (*P. africana*, synonym: *Pygeum africanum* Hook. F) is an evergreen tree species that occurs within the sub-Saharan countries of Africa including Angola, Burundi, Cameroon, the Democratic Republic of Congo, Equatorial Guinea, Ethiopia, Kenya, Lesotho, Madagascar, Malawi, Mozambique, Nigeria, Rwanda, Sao Tome, South Africa, South Sudan, Swaziland, Tanzania, Uganda, Zambia, and Zimbabwe (Komakech et al., 2019). The genus name *Prunus* derives from the Latin word that refers to the plum family and the binomial Latin name *P. africana* indicates the African origin of the species (Komakech et al., 2017). *P. africana* is a highland forest tree, growing in the humid and semi-humid highlands and humid midlands. The wild tree species is found
mainly in tropical forests at altitudes of approximately 900–3,400 m above the sea level, with a mean annual rainfall and temperature of 890–2,600 mm and 18–26°C, respectively (Komakech et al., 2017). The mature *P. africana* tree is approximately 10–25 m high with open branches (Fig. 1a). The outer part of the stem bark is corrugated or rough and black to brown in color (Fig. 1b). The leaves are alternate and simple, approximately 8–20 cm in length, dark green on the top, and pale green at the bottom with mildly serrated margins (Fig. 1c). The flowers are small, white or greenish, hairy, and borne in bunches. The fruits are spherical, purplish-brown, and bilobed, with a thin tough pericarp (Fig. 1d). The seeds are yellowish-brown and oval in shape (Fig. 1e) (Komakech et al., 2017).

- Figure 1 -

*Prunus africana* has been used in the treatment and management of several diseases including benign prostatic hyperplasia, prostate cancer, gonorrhea, chest pain, fevers, gastrointestinal conditions, urinary disorders, malaria, diabetes, obesity, mental illness, hypertension, infertility, and kidney disease (Steenkamp, 2003; Jimu, 2011; Komakech et al., 2019). Owing to its medicinal importance, a micropropagation protocol for *P. africana* was recently developed to meet the ever increasing demand for it (Komakech et al., 2020). However, as an important medicinal plant, providing features that enhances its accurate taxonomic identification and authentication is pivotal (De Souza et al., 2018). Microscopic observation is one of the important approaches to identify characteristic features that could be used to standardize medicinal plant characterization. Although a study focusing on the anatomy of *P. africana* bark and wood structure has been published (Kotina et al., 2016), there is limited information regarding its leaf, fruit, and seed anatomy. This study thus attempted to provide anatomical characteristics of *P. africana* leaf, seed, fruit, and stem which could be important additional features for its accurate plant identification, quality control, and phylogenetic studies in the future.

2. Material And Methods

The Natural Chemotherapeutics Research Institute, Ministry of Health, Uganda provided the *P. africana* sample for the purpose of this study. The voucher specimen number KIOM201901022377 was deposited in the Korean Herbarium of Standard Herbal Resources (Index Herbarium code: KIOM) at the Korea Institute of Oriental Medicine (KIOM), South Korea. The dried leaves and seeds for the anatomy study were stored in water for 24 h, followed by maceration in boiling water for 10 min. An ascending ethanol series (50–100 %) was used to dehydrate the tissues for 1 h at each concentration. The samples were then embedded using Technovit® 7100 (Heraeus-Kulzer, German), based on a previously described protocol (Yeung and Chan, 2015). After complete polymerization, 10 µm sections of the resulting resin blocks were prepared using a microtome (SM 2010R; Leica, Wetzlar, Germany) with a tungsten carbide knife. The resin films containing the tissue sections were attached to the glass slides using warm water, followed by staining with Toluidine Blue O before the mounting using Permount® (Fisher Science, Hampton, USA). The slides of the studied material were observed under a microscope (BX53; Olympus, Tokyo, Japan) and photographed using a digital camera (DP-51; Olympus, Tokyo, Japan).
For the leaf cuticle morphological observations, living material samples were stored using 70 % ethanol, then cut into small pieces (1.0 cm × 1.0 cm). For the light microscopic observations, the samples were dipped in 6 % sodium hypochlorite for 8 h. The samples were then thoroughly washed in distilled water. The epidermis of both surfaces of the leaves was peeled off using a single-edge blade (DN-52, Dorco, Seoul, Korea), colored in 1 % safranin-50 % ethanol for 3 min and mounted in Canada balsam. The mounted slides were examined under a light microscope (Olympus BX-53, Olympus, Tokyo, Japan), and captured using a digital camera (Olympus DP21, Olympus, Tokyo, Japan). The distribution of the epidermal types and stomata density were recorded and compared from the central part of the leaves. The cuticle morphological terminology used to describe the leaves in this study followed previously published indications (Wilkinson, 1979; Evert, 2006).

3. Results And Discussion

Leaf morphology plays an integral role in plant taxonomy, identification, and systematics (Viscosi and Cardini, 2011). In this study, the leaves of \textit{P. africana} were observed to exhibit a smooth surface and hypostomatic nature with stomata size ranging 18.90–(22.34)–26.90 × 15.41–(18.40)–21.22 µm (Table 1). The leaves were also observed to display an isodiametric or irregular cell arrangement (Table 2). Tetracytic and anisocytic stomata complexes and three anticlinal cell wall types (straight, undulated, and straight/curved) were observed on the leaf surfaces (Fig. 2A and B). These stomatal complex types are important factors in accurate plant classification (Abdulrahaman et al., 2009), determination of plant origin, evolution, and phylogenetic relationships (Hong et al., 2018).

### Table 1
Overview of representative stomatal characteristics of \textit{P. africana}. HP-hypostomatic; AB-abaxial surface; Act-actinocytic; Ani-anisocytic; Ano-anomocytic; Tet-tetracytic.

| Prunus species | Stomatal complex | Size of stomata (µm) |
|----------------|------------------|----------------------|
|                | Position | Type | Length | Width |
| \textit{Prunus africana} | HP | AB | Ani, Tet | 18.90-(22.34)-26.90 | 15.41-(18.40)-21.22 |

### Table 2
Overview of representative leaf epidermal surface characteristics of \textit{P. africana}. AD-Adaxial surface; AB-abaxial surface; iso-isodiametric, St-straight; cur-curved; und-undulate; ft-flat; DR-druse-shaped crystal; ST-Star-shaped crystal; SS-short simple trichomes; LS-long simple trichomes; GT-glandular trichomes. -, absent; +, presence; ++, dominant.

| Prunus species | Primary sculpture | Crystals | Trichomes |
|----------------|------------------|----------|-----------|
|                | Outline | Anticlinal wall | Periclinal wall | DR | ST | SS | LS | GT |
| \textit{Prunus africana} | AD | iso | und | ft | - | ++ | - | + | - |
| AB | iso | cur/st, und | ft | - | ++ | - | - | - |
In addition to morphology, leaf anatomy has gained importance as a key tool in plant taxonomy over the years (Araújo et al., 2010; Kolb et al., 2020). In this study, the transverse-section through the leaf mid-vein and petiole of *P. africana* showed the presence of characteristic druse crystals (Fig. 2C, E, and H), calcium oxalate crystals distributed in all plants and known to protect plants against herbivores (Franceschi and Nakata, 2005). The shape of these crystals is reportedly genetically controlled which explains their consistency in a given plant species (Ilarslan et al., 2001). The presence of these druse crystals might thus play an important role in plant taxonomy since they occur in various morphological shapes as per given plant species, including druses, prisms, styloids, raphides, and crystal sand that vary from one plant species to another (Konyar, et al., 2014). Previous studies also reported the presence of druse crystals in the leaves of other *Prunus* species including *P. serotina* (Lersten and Horner, 2006) and *P. virginiana* (Lersten and Horner, 2004).

Our observation showed that the transverse-section of the *P. africana* petiole contained vesicular bundles with prominent xylem vessels arranged in a circular pattern (Fig. 2D and E). The *P. africana* mid-leaf transverse-section revealed the presence of upper and lower epidermal layers with cuticle and a single layer of end-to-end densely packed epidermal cells with large vacuoles (Fig. 2F and G). It is an important structure that protects the plant against moisture loss, microbial, and physical harm (Crang et al., 2018). The palisade layer located on the adaxial side just beneath the upper leaf epidermis was made of closely packed cylindrical cells, and a spongy layer located in the abaxial side of the leaf with loosely arranged irregular shaped cells and wide intercellular space (Fig. 2G and H).

The young stems have green and glabrous surface (Fig. 3A). The anatomical section of the stem showed the presence of a layer of isodiametric cells covered by a smooth cuticle. The cortical parenchyma inner regions contain the intercellular spaces. Sclerenchyma cells were absent. Primary vascular bundles xylem and phloem vessels were arranged in the shape of a ring (Fig. 3B). The pith was centrally located, consisting of spongy parenchyma cells (Fig. 3B–D). The cortex consisted of isodiametric, thin-walled parenchyma cell layers containing druse crystals (Fig. 3D). Druse crystals were observed in the cortex region of the stem. The mature bark is corrugated or rough and black to brown in color. It contains brown dots and/or patches of lenticels and adherent scales (Fig. 3E). The microscopic examination of the stem bark of mature *P. africana* (Fig. 3F) showed mainly the presence of rhytidome-secondary phloem and the oldest periderm densely arranged in multiple layers (Fig. 3G). Furthermore, a cluster of stone cells was observed scattered among the rhytidomes. Sclerenchyma was observed within the phelloderm of mature stems (Fig. 3H).

The structures observed in *P. africana* stem bark including the periderm in multiple layers, intercellular spaces, secondary phloem, the sclerenchyma tissues, occurrence of druse crystals, almost exclusively simple perforation plates, and stone cells are shared by other *Prunus* species that were previously studied including *P. serotina*, *P. Avium*, *P. Pennsylvanica*, and *P. Pennsylvanica* (Bastin, 1895); indicated that they all belong to the genus *Prunus*. 
The anatomy of the *P. africana* pericarp showed the presence of epicarp, mesocarp, and endocarp with a thickness of approximately 350–400, 300–350, and 30–50 µm, respectively, and vascular bundles (Fig. 4A, B, C).

Seed anatomy plays a vital role in the taxonomy of plants (Vaughan, 2009). In this study, the transverse section through the seed of *P. africana* showed the presence of testa with a thickness of approximately 50–60 µm, plumule, and sclenchyma tissue.

### 4. Conclusions

The internal structures of the plants play critical roles in the understanding of the relationships between the taxa. Consequently, the results obtained in this study will play a crucial role in the taxonomy of *P. africana*.

### Declarations

#### Authors contribution

RK conceived the original research plans, collected experimental materials, conducted the experiments, and wrote this manuscript. SY, JHS, and GC carried out anatomical study and authentication of the *P. africana* used in this study. YGK wrote the manuscript. FO and GNK collected the sample, wrote, and revised the manuscript. MGM technically revised the manuscript. YK supervised all the experiments and is the corresponding author.

#### Ethics approval and consent to participate

Not applicable

#### Data availability

The raw data supporting the results of this article will be made available through the corresponding author upon reasonable request.

#### Competing Interest

The authors declare no conflict of interest.

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**Figures**
Figure 1

Botany of Prunus africana. a. P. africana tree. b. Stem of P. africana with part of its bark harvested for medicine purpose. c. Leaf of P. africana. d. Fruit of P. africana. e-f. Seeds of P. Africana
Figure 2

Light microscope micrographs of leaf characteristics of P. africana. A. Leaf adaxial surface. B. Leaf abaxial surface. C. Druse-shaped crystals. D, E: transverse sections of leaf petiole, F-H: transverse sections of leaf blade. C, E, H: Observation was made on DIC mode of light-microscope. ep: epidermis, ics: intercellular space, pa: palisade parenchyma, spo: spongy parenchyma, vb: vascular bundle, xy: xylem, arrows: druse crystal. Scale bars for A-B: 50 μm; C: 20 μm.

Figure 3

The stem morphology and anatomy of Prunus africana. A. Young stem morphology. B: Transverse section of a young stem. C-D: Longitudinal sections of a young stem. E. Mature stem bark morphology. F. Dried stem bark. G-H. Rhytidome. Observation on DIC mode of light-microscope. cor: Cortex. ep: Epidermis, pf: Phloem fiber, ph: Phloem, xy: Xylem. Arrows: Druse crystal. per: Periderm, sc: Sclerenchyma. G and H scale of 200 μm.
Figure 4

Fruit and seed anatomy of Prunus africana. A, B, C: transvers section of pericarp. D: Transvers section of seed. epc: Epicarp, mec: Mesocarp, end: Endocarp, vb: Vascular bundle. cot: cotyledon, pl: Plumule, ts: Testa. Arrows: Sclenchyma.