Comparative stem and petiole anatomy of West African species of Momordica L (Cucurbitaceae)

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Petiole and stem gross anatomy of seven West African species of Momordica L of the family Cucurbitaceae were studied. This was with a view to exploiting their systematic and taxonomic significance; this is because other studies on them were based on morphology. Representatives of the five species were obtained from various parts of the West African sub-region and passed through standard treatments to make permanent anatomical slides for the study. Micrographic evidences of distinguishing and affinity taxonomic features were made. Variations in petiole and stem anatomical attributes were obvious that they could be used as systematic evidence to taxonomically delineate these taxa even at species level. The evidences are used to locate the species in the tribe Cucurbitoideae. The occurrence of grit cells below the epidermis of the petiole of Momordica cabraei is remarkable and separates it from others; this is reported for the first time. Whereas in Momordica cissoides, brachysclerids are found interspersing the cells of the epidermal layer of the petiole which is also reported for the first time. Both stem and petiolar vascular bundle quantitatively separates the species. Cortical parenchyma cellular layers differentiates the species and therefore delineated them. Momordica balsamina stands apart with the possession of stemic 15 to 17 cellular layers of sclerenchymatous tissues. In M. cabraei, the grit cells are present in stem where it is scattered across the tissues whereas in Momordica multiflora, where it was lacking in petiole, packs are arranged below stemic epidermal tissues. The occurrence of starch deposit across various tissue layers of M. multiflora petiole also stands it out. Various other distinguishing features are discussed. The use of petiolar and stemic anatomical features in systematic description of Momordica species is maiden and innovative and reported for the first time.

Key words: Momordica, Cucurbitaceae, stem, petiole, systematics, taxonomy, cucurbitoideae.

INTRODUCTION

Momordica L is of the tribe Joliffia in the family Cucurbitaceae, sub-family Cucurbitoidea (Jeffrey, 1964, 1980). The genus is represented in West Africa by seven species (Hutchinson and Dalziel, 1954). Momordica like other Cucurbits are frost sensitive and therefore, confined to warmer parts of the globe (Tsuchiya and Gupta, 1991, Uno et al., 2001). The genus share cucurbits common characteristics (Purseglove, 1968). Momordica is recognisable in the field with such obvious features as coarse tendrils usually extra-axillary, bearing vines with unisexual flowers often yellowish or white with inferior and leathery berries/pepo (Ndukwu, 1988; Aguoru and Okoli, 2008). The stems are herbaceous, angular hollow, climbing by means of tendrils (Gill, 1988). Momordica species are plants with enormous potential as source of food and drug (Aguoru and Ogaba, 2010); they are associated with varied ethno-botanical uses and occupy special place in the lives and activities of many West African tribes (Aguoru and Okoli, 2008; Burkill, 1985; Okoli, 1984; Dalziel, 1937; Aguoru and Ogaba, 2010). Various studies on Momordica have been based more on morphological features (Aguoru and Okoli, 2008). The
use of plant anatomy (internal structures) in the elucidation of taxonomic and systematic relationships is not new and has been reported (Edoga and Okoli, 1997, 2001; Mbagwu and Edoga, 2006; Nwachukwu and Mbagwu, 2007; Vaughan, 1970; Singh and Dattan, 1980; Ndukuw, 1988; Esau, 1965; Metcalf and Chalk, 1979).

Despite the availability of these studies, no investigation has been conducted on not just the taxonomic usefulness of the stem and petiole anatomy of *Momordica* species but on the entire anatomy of the two organs of the *Momordica* species. This paper therefore reports for the first time the basic anatomical features of the stem and petiole of the West African species of *Momordica* which are also of systematic and taxonomic values.

**MATERIALS AND METHODS**

Representatives of the seven species were obtained from various parts of the West African sub-region and passed through standard treatments to make permanent anatomical slides for the study. Mature fresh and hot water revived dry stems and petioles were used for the study. The materials were fixed in FAA (1:1:8, 40% formaldehyde : glacial acetic acid : 70% ethyl alcohol v/v) for at least 48 h. The tissues which had been so fixed were later used for section preparation following the method of Aguoru and Okoli (2008) with modifications. The stem and petiole tissues were rinsed in several changes of distilled water, placed in glass vials and dehydrated through alcohol series (30, 50, 70, 95 and 100%) for 2 h in each. The dehydrated tissues were infiltrated with wax by passing them through different proportions of alcohol and chloroform (3:1, 1:1, and 1:3) v/v for 3 h in each. As the chloroform gradually replaced, the alcohol, pure chloroform and wax were put in the vials containing the tissues. This was to gradually infiltrate the tissue with wax, which would make them hard enough for microtomy.

The vials were left on a hot plate (37 to 40°C) for 24 h before being transferred to oven (60 to 65°C). This was meant to evaporate the chloroform and facilitate the complete infiltration of the tissues with wax. After a period of 72 h with constant addition of wax, the tissues were embedded in paraffin wax by use of metal moulds and molten wax. This was accomplished by a quick orientation of the plant materials in mould with a hot mounted needle and forceps, and cooling in iced-water later. The moulds were later removed and the “wax-cubed” containing the tissues were trimmed and sectioned using a Shandon micromtome at 15 to 20 µm. The section in wax ribbon was collected on slides already smeared with egg albumen. The slides were placed on hot plate at 40°C for 4 min to enable the ribbons expand and kept in an oven at 40°C till required. The sections were de-waxed in pure xylene and rehydrated in alcohol series in the order of 100, 95, 70, 50 and 30% for a few minutes in each. Staining was achieved by placing drops of 1% safranin on the sections for about 2 min, washed of with water and sections dehydrated through alcohol series starting from 50, 70, 95 and 100% for a few minutes in each. Clearing was done with xylene and mounted in Canada balsam. The slides so prepared were dried on a hot plate at 30°C. A minimum of 30 anatomical sections made from specimens obtained from various different locations were viewed to establish number of cell layers that constitute each tissue section and number of vascular bundles for each species. Photomicrographs were taken using a Leitz Wetzlar Ortholux Microscope fitted with Vivita camera from best anatomical sections.

**RESULTS**

The comparative anatomical features of the petioles and stems of the seven species of *Momordica* found in West Africa are summarised in Table 1 and displayed in the Figures 1 and 2. The petiole of *Momordica charantia* shows a piliferous, single layered epidermis made up of oval shaped cells closely followed by two layers of collenchymatous cells and seven layers of sclerenchyma cells, 10 bicollateral vascular bundles, which appears thicker on the outer portion. The hypodermal cells are oval shaped; the petiole has a pith. *Momordica cissoids* petiole epidermal cells are single layered having grit cells at regular intervals, one to two layers of collenchyma cells and three to four layers of sclerenchyma cells. The vascular bundles are eight in number and there is the presence of pith. *Momordica multilora* petiole has a single layered epidermal cell, one to two layers of collenchymatous cells and four to five layers of sclerenchyma cell with 18 number vascular bundles. There is the presence of pith which is non-septate. *Momordica cabraei* petiole epidermis is single layered, below it occurs a double layered grit cells, four to six layers of sclerenchyma and one to two layers of parenchyma. 10 number bicollateral vascular bundles and two to three layers of grit cells surround the vascular bundles especially around the ridges; pith is absent. *Momordica foetida* petiole epidermis is piliferous and single layered, sclerenchyma cells three to four layers with a single layer of parenchyma which are elongated, with 10 bicollateral vascular bundles and non septate pith. *Momordica angustisepala* petiole has single layered epidermis followed by a single layered collenchymatous cells and four to seven layers of Sclerenchymatous cells having thickened walls. 18 number bicollateral vascular bundles; pith is present. *Momordica basalmina* petiole is winged, covered with single layered epidermis, two to three layers of collenchyma and about three layers of Sclerenchymatous cells. Six number bicollateral vascular bundles; pith is absent. The stem of *Momordica charantia* like the petiole has an oval shape, single cell layered piliferous epidermis, four to five cell layers of collenchyma and five to six layers of sclerenchyma cells, 10 bicollateral vascular bundles and a hollow cylinder. *Momordica cissoides* stem has single cell layered epidermal cells, around the ridges five to six layers of sclerenchyma cells around the furrows directly below the epidermis eight to 10 layers of collenchyma, two to three layers of sclerenchyma and single layer of parenchymatous cells surrounding 12 bicollateral vascular bundles. In *Momordica multilora* stems have furrows and ridges. The epidermis on the ridges are single layered and directly below it is a single layer of brachysclerids (grit cells) six to seven layers of sclerenchyma and same number of layers of parenchyma. On the furrows, epidermal cell layers are two followed by two to three layers of collenchymatous cells before a single layer of sclerenchyma cells. The
Table 1. Summary of the anatomical features of the Petiole and stem of the *Momordica* species studied.

<table>
<thead>
<tr>
<th>Plant part</th>
<th>Features</th>
<th><em>M. charantia</em></th>
<th><em>M. cabrae</em></th>
<th><em>M. cissoides</em></th>
<th><em>M. foetida</em></th>
<th><em>M. multiflora</em></th>
<th><em>M. angustisepala</em></th>
<th><em>M. balsamina</em></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Petiole</strong></td>
<td><strong>Nature of epidermis</strong></td>
<td>Piliferous oval shaped albumen</td>
<td>Single layer</td>
<td>Single layer non-piliferous</td>
<td>Single layer and piliferous</td>
<td>Single layer</td>
<td>Single layer</td>
<td>Single layer</td>
</tr>
<tr>
<td><strong>Sclerenchyma</strong></td>
<td>1-7 layers</td>
<td>4-6 cell layers</td>
<td>3-4 layers</td>
<td>3-4 layers</td>
<td>4-5 layers</td>
<td>4-7 layers</td>
<td>3 layers</td>
<td></td>
</tr>
<tr>
<td><strong>Parenchyma</strong></td>
<td>Absent</td>
<td>1-2 layers</td>
<td>Absent</td>
<td>Elongated 1-2 layers</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
<td></td>
</tr>
<tr>
<td><strong>Collenchyma layer thickness</strong></td>
<td>2 cell layers thick</td>
<td>Absent</td>
<td>1-2 layers thick</td>
<td>Absent</td>
<td>1-2 layers</td>
<td>Single layer</td>
<td>2-3 layers</td>
<td></td>
</tr>
<tr>
<td><strong>Number of vascular bundles</strong></td>
<td>10 bi-collateral vascular bundle</td>
<td>10 bi-collateral vascular bundle</td>
<td>8 bi-collateral vascular bundle</td>
<td>10 bi-collateral vascular bundle</td>
<td>18 bi-collateral vascular bundle</td>
<td>18 bi-collateral vascular bundle</td>
<td>6 bi-collateral vascular bundle</td>
<td></td>
</tr>
<tr>
<td><strong>Other distinguishing features of each species</strong></td>
<td>Pith present with septum</td>
<td>2 grit cell layers under epidermis</td>
<td>Grit cells interspersed epidermis</td>
<td>Pith present</td>
<td>Starch deposits at various tissue layers</td>
<td>Pith present without septum</td>
<td>Winged</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Stem</strong></th>
<th><strong>Nature of epidermis</strong></th>
<th>Piliferous single layer</th>
<th>Single layers cuticularised</th>
<th>Single layer</th>
<th>Single layer</th>
<th>Single layer</th>
<th>Single layer</th>
<th>Single layer and cuticularised</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Collenchyma</strong></td>
<td>4-5 layers</td>
<td>Absent</td>
<td>8-10 layers</td>
<td>2-3 layers</td>
<td>2-3 layers on furrows only</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td><strong>Cortical parenchyma</strong></td>
<td>4 cell layers thick</td>
<td>6-8 layers thick</td>
<td>Single layers</td>
<td>2-3 layers</td>
<td>6-7 layers</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td><strong>Sclerenchyma layer thickness</strong></td>
<td>6 layers thick</td>
<td>6-7 layers thick</td>
<td>2-3 layers</td>
<td>6-7 layers</td>
<td>6-7 layers</td>
<td>8-9 layers</td>
<td>15-17 layers with very thick walls</td>
<td></td>
</tr>
<tr>
<td><strong>Number of vascular bundle</strong></td>
<td>10 bi-collateral vascular bundle</td>
<td>10 bi-collateral vascular bundle</td>
<td>12 bi-collateral vascular bundle</td>
<td>12 bi-collateral vascular bundle</td>
<td>20 vascular bundle with special sieve tubes present</td>
<td>11 bi-collateral vascular bundle</td>
<td>18 bi-collateral vascular bundle with circular cylinder carrying thick walls.</td>
<td></td>
</tr>
<tr>
<td><strong>Other distinguishing features species each studied</strong></td>
<td>Hollow cylinder</td>
<td>Brachysclerids scattered</td>
<td>On the ridges sclerenchyma is 5-6 layers</td>
<td>Pith present</td>
<td>Grit cells directly below epidermis on ridge</td>
<td>Pith present with septum</td>
<td>Circular cylinder walls around the ridges are thickened</td>
<td></td>
</tr>
</tbody>
</table>
Figure 1. Petioles of the various *Momordica* species studied. A, Epidermis; B, cortex; C, vascular bundle; D, grit/stone cells (Brachysclereids).
stem possesses 20 numbers of vascular bundles with special sieve tubes. The stem of *M. cabraei* has a single layer of outer epidermis which is one cell layer thick and covered by a thin cuticle, below the epidermis are two to
three layers of brachysclerids followed by six to seven layers of sclerenchyma cells and six to eight layers of parenchyma, two bicollateral vascular bundle and circular central pith. There is the presence of stellar Sclerenchymatous bundle sheath reinforcement of the vascular bundle. *M. foetida* has a single layered piliferous epidermis, two to three layers of collenchyma, six to seven layers of sclerenchyma and two to three layers of parenchyma, 12 bicollateral vascular bundle; pith is present but with a septum. On the ridges, there are wing like structures which carry three to four circular holes in them and there is a mixture of collenchyma and parenchyma. *M. angustisepala* stem carries a single layer epidermis without cuticle, eight to nine layers of sclerenchyma and 16 bicollateral vascular bundle; the other cell layers are missing; pith is present. The stem of *M. balsamina* is covered with single layer of epidermal cells and cuticularised. 15-17 layers of sclerenchyma cells are with highly thickened walls, and 18 bicollateral vascular bundles with a broad circular cylinder. The cylinders are reinforced by thick walls especially on the ridges.

**DISCUSSION**

Anatomical evidences have been exploited in delimiting taxa (Metcalfe and Chalk, 1979; Aguoru and Okoli, 2008c). The petiole and stem anatomy of the *Momordica* species showed marked variations which are of systematic and taxonomic significance and are reported for the first time. The presence of pith with septum on the petiole of *M. charantia* separates it from the other six species. In *M. cabraei* petiole, two cell layers of grit cells appear below the epidermis whereas in *M. cissoides*, the brachysclerids interspersed the epidermal cells. The petiolar grit cells are lacking in the other five species; whereas this suggests closer affinity between the two species, their positioning separates them but the presence also distinguishes them from the other species. The occurrence of starch deposits in various tissue layers of the petiole of *M. multiflora* stands it out. The number of vascular bundles present in the petiole of the various species ranges from six to 18 numbers, whereas they are 18 in *M. multiflora* and *M. angustisepala*, 10 in *M. charantia*; *M. cabraei* and *M. foetida*, they are eight in *M. cissoides* and six in *M. balsamina* and could be used in phyllogenetic tree construction. The number of layers of hypodermal sclerenchyma confirms that all species could be put in the same genus as it ranges between one to seven layers in all species examined. The presence or absence of hypodermal parenchyma could be used as distinguishing character amongst the species and strengthens the vascular bundle quantitative evidence in phyllogenetic tree construction. Stem anatomical display show that cortical parenchyma could be a distinguishing feature as there is marked variation in layer thickness amongst the various species. The Sclerenchymatous layer of the cortex connotes closer taxonomic ties amongst the species which could be used to enclose them in the same genus except for *M. balsamina* where it distinguished the species from others. Whereas in other species, the layers are between six to nine cell layers, in *M. balsamina* it is between 15 to 17 cell layers thick standing it out taxonomically. Grit cells are present in *M. cabraei* scattered across tissue layers, *M. multiflora* arranged directly below epidermis and lacking in *M. cissoides* where it occurred in petiole. The nature of epidermal cells suggests taxonomic affinity amongst the species. The present result indicates and strengthens the fact that the species could continue to be treated as same genus but separate as they are as species. It also affirms that the genus belongs to the tribe Cucurbitoideae.

An indented taxonomic (diagnostic) key based on stem and petiole anatomical characters of the *Momordica* L species studied in the present work is presented below.

1. Grit/stone cells/sclereids present in either or both stem and petiole anatomical section.

2. Grit/stone cells found only in petiole section

3. Grit/stone cells two layers below the epidermis of petiole section, petiole with 10 bicollateral vascular bundles

4. Both cortical parenchyma and collenchyma absent in stem anatomical section with variations in number of vascular bundles and sclerenchyma thickness.

5. Bicollateral vascular bundle 11 with eight to nine layers of sclerenchyma, pith with septum

5' bicollateral vascular bundle 18 with 15 to 17 layers of sclerenchyma made of cells with thick walls around the ridges

6. Six sclerenchyma layers

7. 10 bicollateral vascular bundle, stem section having
hollow cylinder, pith absent

\[ M. \textit{charantia} \]

7’ 12 bicollateral vascular bundle, pith present

\[ M. \textit{foetida} \]

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