

Morphological and molecular barcode analysis of the medicinal tree *Mimusops coriacea* (A.D.C). Miq. collected in Ecuador

Katherine Bustamante¹, Efrén Santos-Ordóñez^{Corresp., 2,3}, Migdalia Miranda⁴, Ricardo Pacheco², Yamilet Guitiérrez⁵, Ramón Scull⁵

1 Manuscript Title

5 Authors

6 Katherine Bustamante¹, Efrén Santos-Ordóñez^{2,3}, Migdalia Miranda⁴, Ricardo Pacheco-Coello², Yamilet Gutiérrez⁵, Ramón Scull⁵.

¹Universidad de Guayaquil. Facultad de Ciencias Químicas. Ciudadela Universitaria “Salvador Allende”. Ave. Kennedy S/N y Av. Delta. Guayaquil. Ecuador telef. 593- 229 3680/2293379.

11

12 ²ESPOL Polytechnic University, Escuela Superior Politécnica del Litoral, ESPOL, Centro de Investigaciones Biotecnológicas del Ecuador, Campus Gustavo Galindo, Km. 30.5 vía Perimetral, P.O. Box 09-01-5863, Guayaquil, Ecuador

15

16 ³ESPOL Polytechnic University, Escuela Superior Politécnica del Litoral, ESPOL, Facultad de Ciencias de la Vida, Campus Gustavo Galindo, Km. 30.5 vía Perimetral, P.O. Box 09-01-5863 Guayaquil, Ecuador.

19

20 ⁴ESPOL Polytechnic University, Escuela Superior Politécnica del Litoral, ESPOL, Facultad de

21 Ciencias Naturales y Matemáticas. Campus Gustavo Galindo. Km 30.5 vía Perimetral.

22 Guayaquil. Ecuador. Email.

23

24 ⁵Instituto de Farmacia y Alimentos. Universidad de la Habana. 222 y Ave 23. La Coronela. La Lisa. Ciudad Habana. Cuba.

26

27 Corresponding Author:

28 Efrén Santos-Ordóñez^{2,3}

29 ²ESPOL Polytechnic University, Escuela Superior Politécnica del Litoral, ESPOL, Centro de Investigaciones Biotecnológicas del Ecuador, Campus Gustavo Galindo, Km. 30.5 vía Perimetral, P.O. Box 09-01-5863, Guayaquil, Ecuador

30 ³ESPOL Polytechnic University, Escuela Superior Politécnica del Litoral, ESPOL,
Facultad de Ciencias de la Vida, Campus Gustavo Galindo, Km. 30.5 vía Perimetral,
P.O. Box 09-01-5863 Guayaquil, Ecuador.

31 Email address: gsantos@espol.edu.ec

36

37 **Abstract (limit to 200 words maximum? See journal guidelines)**

38 Background

39 *Mimusops coriacea* (A.D.C). Miq., ~~is~~ a species from the Sapotaceae Family,
originated from Africa. ~~*Mimusops coriacea*~~ plants were introduced to coastal areas in
Ecuador, ~~and with tissues material from the tree~~ ~~are now regularly~~ extensively used as
a traditional medicine to treat various human diseases in ~~humans~~ Ecuador. Different
therapeutically uses of the species include: analgesic, inflammation and pain
~~purposes—relieve associated with~~ bones and articulation-related diseases.
Furthermore, ~~tissues from~~ *M. coriacea* could be used as an anti-oxidant agent.
However, ~~limited research has been focused only in few *Mimusops* species including~~
~~*M. elengino*~~ botanical, chemical, or molecular barcode information related to this
much used species is available. ~~Therefore, botanical, chemical, and molecular~~
~~barcode studies for *M. coriacea* are null.~~ In this study, morphological characterization
was performed in different plant tissues including leaves, stem and seeds ~~from fruits~~.
Furthermore, genetic characterization was performed using molecular barcodes for
rbcL, *matK*, ITS1 and ITS2, using DNA extracted from leaves.

49

50 Methods

51 Macro-morphological description was performed in fresh plant ~~tissues—material~~
including leaves, stem and seeds. For ~~micro-morphological~~ anatomical evaluation,
tissues were embedded in paraffin and transversal dissections were done following
incubation with sodium hypochlorite and safranin for coloration and fixated later in
glycerinated gelatin. DNA extraction was performed using a modified CTAB protocol
from leaf tissues and amplification by PCR was accomplished for the molecular
barcodes *rbcL*, *matK*, ITS1 and ITS2. Sequence and phylogenetic analyses were
performed using blastn and MEGA, respectively, with sequences in the GenBank.

58

59 Results

60 ~~For the first time, morphological and genetic characteristics were performed in the *M. coriacea* (A.D.C). Miq. Detailed morphological characteristics were obtained in the different tissues analyzed. Indicate the most significant macro- and micro morphological characterization results obtained. Sequence analysis revealed that amplicons were generated using the four barcodes selected. Phylogenetic analysis revealed-indicated that the barcodes *rbcL* and *matK*, were not discriminated between species, and different genus were grouped in one clade of the subfamily Sapotoideae. On the other hand, the ITS1 and ITS2 were discriminative at the level of genus and species of the Subfamily Sapotoideae.~~

Commented [PM1]: Just state result

67

68 **Introduction (this Introduction is too short for a full paper (which this paper is) – it needs significant expansion in terms Medicinal properties, Industrial applications and pharmacological properties – data is presented too condensed.)**

69 Sapotaceae is a botanical Family which contains 58 genera of plants with 1271 species. The most important genera from this Family includes *Pouteria*, *Palaquium*, *Madhuca*, *Manilkara*, *Sideroxylon*, *Chrysophyllum* and *Mimusops*. Genera relevant for geographic distribution and pharmacological applications are *Manilkara* and *Mimusops*. In the genus *Mimusops* (Sapotaceae), a total of 45 species have been described and that are distributed in Asia, Africa, Australasia and Oceania. In Ecuador *Mimusops coriacea* (ADC) Miq. is found ... [Give geographical distribution of this species globally and in Ecuador (indicate which coastal regions), as well as in which vegetation types it occurs]. *Mimusops coriacea* (Give a short diagnostic botanical description of this species as found in Ecuador)-

Formatted: Font: Not Italic

Formatted: Font: Not Italic

Formatted: Font: Italic

70 [is a tree reaching a height of 25 meters, with a dense cope and an irregular short trunk, which exhibit and a cracked bark structure. The tree contains simple leaves which that are alternated and clustered with a brilliant color green. The calyx of the flowers contains eight triangular sepals with brown external hairs. – see if this relates to *M. coriacea*]

71 In Ecuador *M. coriacea* This plant species is used in different ethnic groups for medicinal and industrial purposes For example, (Semenya, Potgieter, & Erasmus et al. (2012) indicated that the species is used by the ?ethnic group for ... (significantly expand this paragraph to indicate *M. coriacea* wide ranging medicinal use across different cultural groups in Ecuador and globally)

72 Chivandi et al. (2016) noted the species industrial application by pointing out it use in For similar species, triterpenoids have been described including phenolies and flavonoids (Chanda et al. 2010; Fayek et al. 2012; Kaneria et al. 2012; Baki et al. 2016). (significantly expand this paragraph to indicate *M. coriacea* wide ranging industrial use internationally and in Ecuador)

Commented [PM2]: No value - paragraph focus exclusively on *M. coriacea*

Formatted: Normal, No bullets or numbering, Tab stops: Not at 1.37 cm + 1.38 cm

69

For the genus *Mimusops*, different pharmacological properties have been indicated (rather focus on *M. coriacea*! –if there is no or only limited pharmacological information related to this species (globally or in Ecuador), then indicate so.) including antioxidant, hypoglycemic, and antimicrobial activities (Said et al. 2003; Ali et al. 2008; Baliga et al.

2011;Gami, Pathak, & Parabia2012; Kar et al. 2012; Kiran Kumar et al. 2014; Saradha Gillani et al. 2017 –Place these sources next to facts INSIDE the sentence – not all at end of sentence, which would then imply that they all stated the same info – in which case why cite 8 when 1 will do?)). (Significantly expand this paragraph to indicate *M. coriacea* wide ranging pharmacological properties linked to its medicinal use)

In Ecuador, limited studies have been conducted ~~to on~~ *Mimusops*, being restricted to ... (indicate areas of research and authors)~~species of this genus of the Sapotaceae family;~~ *Mimusops coriacea* is a very important medicinal species in Ecuador, however, little is known about its ~~therefore, this report includes~~ morphological and anatomical features of its leaves, stems and seeds, as well as its~~and~~ molecular barcode. This study investigated these features. Results obtained will be of value ... [indicate value of results]~~studies of the specie~~ *Mimusops coriacea* (ADC) Miq. including leaves, stems and seeds.

87

88 Materials ~~&~~ and Methods

89 Study area description

90 Plant material were collected during [indicate month and year] in “Jardín Botánico”, a protected natural vegetative area located in the North zone of “Las Orquídeas” area next to the Ave. Francisco de Orellana, in the hills of “Cerro Colorado” of Guayaquil city, Guayas Province, Ecuador (coordinates 02°12'13.6800"S 079°53'50.6400"W). Study site description is necessary for terrain features [In physical geography, terrain is the lay of the land. This is usually expressed in terms of the elevation, slope, and orientation of terrain features. Terrain affects surface water flow, which in turn affect the morphology and anatomy, as well as distribution], associated vegetation types, Climate data, and soil information – these have a direct bearing on the presentation of morphological, anatomical and DNA features.

91

8992 Morphological analysis

9093 Plant tissues were ~~recollected in a protected natural vegetative area named “Jardín Botánico” located in the North zone of “Las Orquídeas” area next to the Ave. Francisco de Orellana, in the hills of “Cerro Colorado” of Guayaquil city, Guayas Province, Ecuador (coordinates 02°12'13.6800"S 079°53'50.6400"W).~~ [indicate how many trees were sampled]Samples were collected ~~corresponded to~~from adult plants of approximately 30 m ~~of in~~ height, with ~~the presence of~~ flowers and fruits. Collected plant samples were ~~cured~~ placed at the GUAY herbarium ~~of the Faculty of Natural Sciences~~ of Guayaquil University, with the accession number 13111 (are all specimens isotypes? – seeing that there is only one number - 13111).

9494 ~~Macro-M~~morphological description of different organs was performed on fresh ~~specimentissues~~and mature leaves, stems and seeds with a stereoscope (model: Zeiss LUMAR.V12–(Germany) with a light source MC 1500 and KL 2500 LCF with a power supply Zeiss HBO100, adapted with an ACXION MRc5 camera. ~~The software used was~~ AXION VISION Rel 4.8 (Zeiss, Germany) software was used in; accordance~~ing~~ to the methodology ~~described of~~ (Miranda ~~&~~ and Cuéllar (2000) to analyze ~~.Different characteristics were described in the leaf (n=?) including~~ shape, edge, apex, base, petiole, venation, consistency, and color. Size was measured in length and width of 100 leaves with a micrometer. For the stems, the characteristics

analyzed included shape, color, external and internal surfaces, and fracture. For fruit characterization, 60 fruits and extracted seeds were analyzed in shape and dimensions, seed coat, and endosperm.

For histological analysis, transversal cuts of fresh leaves were performed manually, which were hydrated and clarified with 1% sodium hypochlorite. Tissues were colored with 1% safranin in water, following fixation with glycerinated gelatin according to Gattuso & Gattuso (1999). To analyze anatomical aspects of the leaf epidermis, a longitudinal cut followed with a diaphanization technique was performed. Cleared leaves were obtained with sodium hypochlorite following incubation with 1% safranin in water. Micro-morphological characteristics of cortex were performed to the drug in powder, performing histochemical reactions including: starch determination (Lugol reagent), lignine (1% safranin in water), and essential oil (5% Sudan III solution in 70% ethanol) (Gattuso & Gattuso 1999). Micromorphology of seeds was performed using dried fragmented material following the procedure described above for leaves and cortex.

9396

9497 DNA extraction and PCR:

Leaves from collected samples were ground using liquid nitrogen in the grinder MM400 (Retsch) and stored at -80°C upon DNA extraction. Approximately, 100 mg of leaf was used for DNA extraction using a CTAB protocol with some modifications (Pacheco Coello et al. 2017). PCR was performed using the 2x GoTaq® master mix (Cat. # M7123, Promega) using 0.5 µM of each primer (Table 1). The final volume was 50 µl per reaction. PCR conditions were 95°C to start denaturation; 35 cycles of: 95°C for 30 s, 60°C (for *rbcL*) or 56°C (for *matK*, ITS1 and ITS2) for 30 s, 72°C for 90 s, with a final extension of 72°C for 5 min. Five microliter of PCR reaction was loaded on a 1.5% gel to check for the presence of amplicons. The remaining 45 µl were purified using the Wizard SV Gel and PCR Clean-Up System (Cat. # A9282, Promega) and sequenced commercially (Macrogen, Maryland, USA).

9699

97100 Bio-informatics analysis of sequences

Sequences were trimmed from low quality using FinchTV or Chromas 2.6.5 (Technelysium). Processed sequences were blast (Zhang et al. 2000) in the GenBank using the nucleotide database. Selected sequences with similarities were selected for phylogenetic analysis using MEGA 7.0.26 (Kumar et al., Stecher, & Tamura 2016). For each barcode, the recommended model from the MEGA7 was used for the phylogenetic analysis after alignment with MUSCLE. For the phylogenetic analysis, around 50 accessions for each barcode of different *genera* from the subfamily Sapotoideae were queried from the GenBank (5th December 2018) and the phylogenetic analysis was also performed independently from selected accession from the blast result.

99102

Results

Morphological evaluation of the leaves (all referenced statements MUST be removed from this section and relocated to the Discussion section – only own present results):

The macro-morphological evaluation allowed the observation of oblong leaves of

coriaceous-waxy texture, short petiole, retuse apex, entire border and obtuse base (-Fig. 1) (Miranda & Cuéllar 2000; Gami, Pathak, & Parabia 2012). The venation is a closed type, which corresponds to a reticular system (the veins branch and anastomose with each other forming a network that facilitates the diffusion of liquids); which is very common in the dicotyledons. In this case, of the penninervia type, the vascular system is one of the most advanced systems that ensures nutrition to all parts of the leaf (Gami, Pathak, & Parabia 2012). -relocate to discussion] Macroscopic details of the leaves are shown (Fig. 1). In respect to the dimensions of the leaves (n=?), the mean average value observed for the length of the leaves was 13.56 ± 1.46 cm, and with a width of 7.49 ± 0.65 cm for the width.

Morphological evaluation of the crust: The crust presented a rugose cuticle of intense gray color, with an underneath slightly brown outer surface (Fig. 2A) with rough streaks. The internal surface was reddish brown, fibrous and furrowed (Fig. 2B).

Morphological evaluation of the seeds: In the macro-morphological study, the length and width of the green and ripe fruits, the seeds with the husk and the endosperm of the seeds were taken into account (Fig. 3). The fruit is rounded, contains one or two seeds, with dimensions of 2.97 ± 0.18 cm long and 3.14 ± 0.25 cm wide when green, reducing their size at maturity to 2.89 ± 0.2 cm in length and 2.97 ± 0.25 cm wide. The seeds with a peel are dark brown with 1.66 ± 0.13 cm long by 1.15 ± 0.21 cm wide when the fruit is green and $1.79 \pm 0.09 \times 1.20 \pm 0.09$ cm, when the fruit is ripe, with an increase in size when the fruit ripens.

153 Anatomical evaluation:

Leaves: In the leaf anatomy at the level of a cross section of the central nerve (Fig. 4A) the adaxial surface is convex, slightly wavy and the abaxial face is concave. An enlarged view of the nerve (Fig. 4B) shows a cuticle of waxy texture that covers the entire leaf, and well visible in the macro-morphological study, followed by the epidermis, which is made up of tabular cells, which gives way to the set of cells that form the spongy parenchyma, given the intercellular spaces which are defined. Possible crystals of calcium oxalate are also observed. Bordering the central part of the central nerve, a cord is observed (Fig. 4C) with color red, corresponding to the endodermis, the structure that surrounds the pericycle. In the middle, the conductive tissue formed by the vascular system xylem and phloem is observed (Fig. 4C).

An image of the leaf mesophyll (Fig. 4D) shows a somewhat thick cuticle on the abaxial surface, followed by the epidermis, a parenchyma palisade with elongated cells that at times become stratified. In the same way, the entire center of the structure occupied by the spongy parenchyma is observed, which borders on the upper epidermis that ends with the cuticle, previously mentioned.

The diafanization of a portion of the leaf by the adaxial side showed an epidermis with cells of variable shape and size (Fig. 4E). However, the abaxial epidermis evidenced many anomocytic type stomata, where the epidermal cells surrounding the pair of occlusive cells are not morphologically different from the rest of the epidermal cells (Fig. 4F). A stain with Sudan III reagent at the level of the epidermis, allowed the visualization of bags with essential oils, which took reddish coloration (Fig. 4G).

The information referenced in the literature regarding the characteristics of the leaves is scarce; thus, comparisons with respect to two species of the genus was performed. For *Mimusops elengi* L. (Gami, Pathak & Parabia 2012), authors reported that the leaves are elliptical in shape, little acuminate at the apex, glabrous with an acute base, and petioles 1.3 - 2.5 cm in length. The dimensions of the leaves range between 6.3 - 10.0 cm long by 3.2 - 5.0 cm wide, while *Mimusops hexandra* Roxb (without *Manilkara hexandra* Roxb), presents oblong leaves, rounded at the apex, glabrous, dark green in the beam and clear on the underside, with a dimension of 2.5 - 11 cm long and 1.0 - 6.0 cm wide (Chanda, Nagani & Parekh 2010). Some species genetically similar to the species under study, present some differences especially in the dimensions of the leaves with respect to those studied, which are superior. The microscopic analysis of the powder drug showed different fibers and vascular bundles, in this case belonging to the xylematic tissue, classified as scalariform. Figure 5 shows the observed microscopic characteristics.

Bark: The micro-morphological analysis of the powder drug showed different fibers and the vascular system, belonging to the xylematic tissue, responsible for the transport of the crude sap to the photosynthetic centers and the circulation of the highest percentage of water. The xylematic vessels are classified as scalariform (Fig. 5). Related to the cortex, no referenced information was found.

Seeds: The micro-morphological analysis of the seed powder (Fig. 5), allowed the visualization of a section of the epispem (outer layer of the seed or testa) where the presence of cells of the sclerenchyma tissue corresponding to the supporting tissue is observed. These cells have a well-defined compact arrangement and the walls are slightly thick. The sclerides of the macro-sclerosis type and elements of the conductive tissue was observed. Histochemical reactions on the samples, demonstrated a well-defined red-colored oil pocket that could be observed through the reaction with the Sudan III reagent. Starch granules of ovoid shape and blackish color were also observed when using the Lugol reagent.

For the seeds, significant differences were observed between the evaluated parameters of the whole fruits and their seeds at maturity (Gopalkrishnan, & Shimpi 2011); for *M. elengi* seed husk was light brown to blackish, with measures of 1.7-1.9 cm long and 1.2-1.5 cm wide, which differs from those obtained for the species studied. The endosperm presented dimensions of 1.42 x 1.00 cm when it came from green fruits and 1.43 x 0.91 cm when it came from ripe fruits, decreasing its thickness in this case.

Molecular barcode of *M. coriacea*.

As a complement analysis for characterization and identification of the *M. coriacea* specimen, PCR for the amplification of the loci *rbcL*, *matK*, ITS1 and ITS2 was performed. Amplicons were detected for all the molecular barcodes and the two samples tested (Fig. 1). Accession number of the sequences in the GneBank are indicated (Table 2).

After alignment of the barcodes sequences from the GenBank with the *M. coriacea* specimen, the best model for phylogenetic analysis are shown (Table 3). The phylogenetic analysis revealed that for the barcodes *rbcL* and *matK*, most of the *Mimusops* spp. are clustered together with other genera (Fig. S1). On the other hand, the ITS1 and ITS2 sequences revealed several clades for the different genera including the *Mimusops* (Fig. S2).

Discussion

1. Shape discussion along the same headings (see below) as used in the results section – this is for easier orientation by author and readers
2. Format discussion in terms of what result means in terms of its medicinal use – as indicated in the Title (...of the medicinal tree ...) of this paper.

Morphological evaluation of the leaves

(Miranda & Cuéllar 2000; Gami, Pathak, &Parabia 2012). The venation is a closed type, which corresponds to a reticular system (the veins branch and anastomose with each other forming a network that facilitates the diffusion of liquids); which is very common in the dicotyledons. In this case, of the penninervia type, the vascular system is one of the most advanced systems that ensures nutrition to all parts of the leaf (Gami, Pathak, &Parabia 2012).

Morphological evaluation of the crust

Morphological evaluation of the seeds

Anatomical evaluation

Leaves:

Bark:

Seeds:

Molecular barcode

Analysis of the molecular barcodes is the first step in the characterization of the *Mimulus* spp. for medicinal application. Molecular barcodes are useful to genetically characterized organisms; and different *loci* have been proposed to be universal for land plants (CBOL Plant Working Group 2009). Although, the two proposed *loci* for barcodes are from chloroplast genome and includes the *rbcL* and *matK*, other *loci* including ITS1 and ITS2 are widely used for medicinal plants (Kim et al. 2016). Furthermore, the ITS2 region is suggested as a barcode for species identification over *rbcL* and *matK* (Tehen et al 2014; Zhang et al 2016 - Place these 2 sources next to facts INSIDE the sentence – not all at end of sentence, which would then imply that they all stated the same info – in which case why cite 2 when 1 will do?)).

- 153 The identification of plant material used as a phytotherapeutic product is a challenge in natural products. One of the many drawbacks is the management of vulgar or regional plant names, the lack of knowledge of the organ or the part of the plant where the active ingredients are found, and the recognition of the macroscopic and microscopic characteristics of plant drugs. Macro morphological and micro morphological studies are essential in the control of the quality of plant drugs, as well as significant details to confirm the identity of the plant, and identification of possible adulterants. Therefore, analysis of the morphology and the molecular barcodes is the first step in the characterization of the *Mimulus* spp. for medicinal application. Molecular barcodes are useful to genetically characterized organisms; and different *loci* have been proposed to be universal for land plants (CBOL Plant Working Group 2009). Although, the two proposed *loci* for barcodes are from chloroplast genome and includes the *rbcL* and *matK*, other *loci* including ITS1 and ITS2 are widely used for medicinal plants (Kim et al. 2016). Furthermore, the ITS2 region is suggested as a barcode for species identification over *rbcL* and *matK* (Tehen et al 2014; Zhang et al 2016).

154

155 Conclusions

156 For the first time, the macro and micro-morphological characteristics of the leaves,
stems and seeds, of the *M. coriacea* collected in Ecuador were performed. The
evaluation of the identity of the species, which is classified taxonomically as
Mimusops sp., is a novelty of this work, and was confirmed by using molecular
barcodes. Most important, the ITS1 and ITS2 indicated more resolution at the species
level (*M. coriacea*) than the *rbcL* and *matK*, confirming published results in
medicinal plants. However, further molecular barcode characterization should be
performed in *Mimusops* spp. to further validate resolution at the species level as a
complement for proper identification using morphological characteristics. Further
pharmacognostic analysis will be performed to study medicinal properties of *M.*
coriacea.

157

158 Acknowledgements

159 Identification of samples by the GUAY herbarium of the Faculty of Natural Sciences
of the

160 Guayaquil University is acknowledged.

161

162 References

- 163 Ali AMA, Mozid MA, Yeasmin S, Khan AM, Sayeed MA. 2008. An evaluation of
antimicrobial
164 activities of *Mimusops* Linn. Research Journal of Agriculture and
Biological
270 Sciences 4:871–874.
- 271 Baliga MS, Pai RJ, Bhat HP, Palatty PL, Bloor R. 2011. Chemistry and medicinal
properties of
272 the Bakul (*Mimusops* Linn): a review. Food Research International.
44:1823–
273 1829.
- 274 Baky MH, Kamal AM, Elgindi MR, Haggag EG. 2016. A Review on Phenolic
Compounds from
275 Family Sapotaceae. *Journal of Pharmacognosy and Phytochemistry* 5(2):280–
287.
- 276 CBOL Plant Working Group. 2009. A DNA barcode for land plants. PNAS
106(31):12794–
277 12797.

- 278 Chanda S, Nagani K, Parekh J. 2010. Assessment of Quality of Manilkara Hexandra (Roxb.)
- 279 Dubard Leaf (Sapotaceae): Pharmacognostical and Physicochemical Profile.
- 280 *Pharmacognosy Journal*. 2(13):520–524. DOI:10.1016/S0975-3575(10)80054-9
- 281 Chanda SV, Nagani KV. 2010. Antioxidant Capacity of Manilkara zapota L. Leaves Extracts
- 282 Evaluated by Four in Vitro Methods. *Journal of Biological Sciences* 8(10): 260–266.
- 283 DOI: [10.7537/marsnsj081010.21](https://doi.org/10.7537/marsnsj081010.21)
- 284 Chivandi E, Mukonowenzou N, Berliner D. 2016. The Coastal Red-Milkwood (Mimusops Caffra)
- 285 Seed: Proximate, Mineral, Amino Acid and Fatty Acid Composition. *South African*
- 286 *Journal of Botany* 102: 137–141 DOI: [10.1016/j.jep.2012.03.008](https://doi.org/10.1016/j.jep.2012.03.008)
- 287 Costion C, Ford A, Cross H, Crayn D, Harrington M, Lowe A. 2011. Plant DNA barcodes can
- 288 accurately estimate species richness in poorly known floras. PLoS One. 6(11):e26841.
- 289 DOI: 10.1371/journal.pone.0026841.
- 290 Fayek NM, Monem AR, Mossa MY, Meselhy MR, Shazly AH. 2012. Chemical and Biological
- 291 Study of Manilkara Zapota (L.) Van Royen Leaves (Sapotaceae) Cultivated in Egypt.
- 292 *Pharmacognosy Research* 4 (2):85-91. DOI: 10.4103/0974-8490.94723.
- 293 Gami B, Pathak S, Parabia M. 2012. Ethnobotanical, Phytochemical and Pharmacological
- 294 Review of Mimusops Elengi Linn. *Asian Pacific Journal of Tropical Biomedicine*
- 295 2(9):743–48 DOI:10.1016/S2221-1691(12)60221-4.
- 296 Gattuso MA, Gattuso SJ. 1999. Manual de procedimientos para el análisis de drogas en polvo.
- 297 Editorial de la Universidad Nacional de Rosario Urquiza. Argentina.
- 298 Gillani SS, Shahwar D. 2017. Investigation of Antioxidant Activity in *Mimusops elengi*. J Plant
- 299 BiochemPhysiol 5:202. DOI:10.4172/2329-9029.1000202.

- 300 Gopalkrishnan B, Shimpi SN. 2011. Seeds of *Mimusops Elengi* Linn.
Pharmacognosy and
- 301 Phytochemical Studies. *International Journal of Pharmacognosy and*
Phytochemical
- 302 *Research*. 3(1):13–17.
- 303 Kaneria M, Chanda S. 2012. Evaluation of Antioxidant and Antimicrobial Properties
of
- 304 *Manilkara zapota* L. (Chiku) Leaves by Sequential Soxhlet Extraction
Method. *Asian*
- 305 *Pacific Journal of Tropical Biomedicine* 2 (3 SUPPL.): S1526–1533. DOI:
- 306 [10.1016/S2221-1691\(12\)60448-1](https://doi.org/10.1016/S2221-1691(12)60448-1)
- 307 Kar B, Kumar RBS, Karmakar I, Dola N, Bala A, Mazumder UK, Hadar PK. 2012.
Antioxidant

308 and in Vitro Anti-Inflammatory Activities of Mimusops Elengi Leaves. *Asian*
309 *Pacific Journal of Tropical Biomedicine* (2 SUPPL.): S976–80. DOI:10.1016/S2221-
310 1691(12)60346-3.

311 Kim WJ, Ji Y, Choi G, Kang YM, Yang S, Moon BC. 2016. Molecular identi cation
312 and phylogenetic analysis of important medicinal plant species in genus
313 *Paenoniabasedon* rDNA-ITS, *matK*, and *rbcL* DNA barcode sequences. *Genetics and*
314 *Molecular Research* 15(3). DOI: 10.4238/gmr.15038472gmr.15038472.

315 Kiran Kumar HA, Mandal BK, Mohan Kumar K, Maddinedi Sb, Sai Kumar T,
316 Madhiyazhagan P, Ghosh AR. 2014. Antimicrobial and Antioxidant Activities of Mimusops
317 ElengiSeed Extract Mediated Isotropic Silver Nanoparticles. *Spectrochimica Acta - Part*
318 *A: Molecular and Biomolecular Spectroscopy* 130:13–18.
DOI:10.1016/j.saa.2014.03.024.

319 Kumar S, Stecher G, Tamura K. 2016. MEGA7: Molecular Evolutionary Genetics
320 Analysis version 7.0 for bigger datasets. *Molecular Biology and Evolution* 33:1870-
1874.

321 Miranda MM, Cuéllar AC 2000. Manual de prácticas de laboratorio. Farmacognosia
322 y productos naturales. Ciudad Habana 25-49, 74-79.

323 Pacheco Coello R., Pestana Justo J., Factos Mendoza A., Santos Ordoñez E. 2017.
324 Comparison of three DNA extraction methods for the detection and quantification of GMO
325 in Ecuadorian manufactured food. *BMC Research Notes* 10:758
DOI:[10.1186/s13104-017-](https://doi.org/10.1186/s13104-017-3083-x)
326 [3083-x](https://doi.org/10.1186/s13104-017-3083-x).

327 Saradha S, Ruckmani A, Chokkalingam M, Maignanakumar R, Arunkumar R,
328 Madhavi E, Lakshmi Prabhhu R. 2014. Hypoglycaemic activity of aqueous and

- ethanolic extracts
- 329 of Manilkarazapota seeds in streptozotocin induced diabetic rats. *Int J*
PharmPharmSci
- 330 6(2):434-437
- 331 Schultz J, Gerlach D, Muller T, Wolf M. 2005. A common core of secondary
 structure of the
- 332 internal transcribed spacer 2 (ITS2) throughout the Eukaryota. *RNA* 11: 361–
 364.
- 333 Semenya S, Potgieter M, Erasmus L. 2012. Ethnobotanical Survey of Medicinal
 Plants Used by
- 334 Bapedi Healers to Treat Diabetes Mellitus in the Limpopo Province, South
 Africa.
- 335 *Journal of Ethnopharmacology* 141:(1) 440–45 DOI:
[10.1016/j.jep.2012.03.008](https://doi.org/10.1016/j.jep.2012.03.008)
- 336 Shah PJ, Gandhi MS, Shah MB, Goswami SS, Santani D. 2003. Study of
Mimusops bark
- 337 in experimental gastric ulcers. *Journal of Ethnopharmacology* 89:305–311.

338 Techen N, Parveen I, Pan Z, Khan IA. 2014. DNA barcoding of medicinal plant
 339 material for identification. Curr. Opin. Biotechnol. 25:103–110.

340 Technelysium. Available at <https://www.technelysium.com.au> (accessed 2 October
 2018)

341 Zhang D, Jiang B, Duan L, Zhou N. 2016. Internal transcribed spacer (ITS), an ideal
 342 DNA barcode for species discrimination in *Crawfordia* Wall. (Gentianaceae).
African journal of traditional, complementary, and alternative medicines: 13(6):101-106.
 343 DOI:10.21010/ajtcam.v13i6.15

344 Zhang Z, Schwartz S, Wagner L, Miller W. 2000. A greedy algorithm for aligning
 345 DNA sequences. J Comput Biol 7(1-2):203-14. Available at
 346 <https://blast.ncbi.nlm.nih.gov/Blast.cgi>

348
 349

350 Figure 1. Macro morphological details of leaf from *M. coriacea*.

351 A: retuse apex, B: whole edge, C: obtuse base, D, E and F: closed rib
 352
 353

354 Figure 2. Macro morphological details of crust from *M. coriacea*.

355 A: external surface, B: internal surface
 356
 357
 358

359 Figure 3. Macro morphological characters of fruits and seeds from *M. coriacea*.

360 a) green fruit, b) ripe fruit, c) seeds green fruits with peel, d) seeds ripe fruits with
 peel,
 361 e) endosperm green seeds, f) endosperm mature seeds

362
 363

364 Figure 4. Microscopic characteristics of leaf from *M. coriacea*.

365 Transversal section of the central nerve of the leaf (I): A, central nerve of the leaf;
B and C,

366 enlarged view of the central nerve; D, mesophilic; Cu, cuticle; Ep, epidermis; COC,
calcium

367 oxalate crystals; SP, spongy parenchyma; VS, vascular system; En, endodermis; AdE,
 368 adaxial
 369 epidermis; PP, palisadeparenchyma; AbE, abaxial epidermis.
 370 **Diafanized of the leaf (II):** E, adaxial epidermis; F and G, abaxial epidermis; EpC,
 371 epidermal
 372 cells; S, stomata; EO, essential oils.

373 **Figure 5. Powder drug characteristics of *M. coriacea*.**

374 VS: vascular system, F: fibers, S: starch, ST: suberoustissue, SF: septate fibers,
 375 COC: calcium oxalate crystal, SC:sclerides cells, MS: macrosclerides, OB: oil bag,
 376 SG: starch granules

377
 378
 379 **Figure 6. Gel electrophoresis of amplicons generated for the molecular barcodes
 with the**

380 **genomic DNA of *M. coriacea*.** **A** amplification of rbcLA_F/ rbcLA_R. **B**
 Amplification of
 381 matK_3F_KIM f/matK_1R_KIM R. **C** amplification of ITS 5a_F/ITS 4_R (ITS I). **D**
 382 amplification of S2f/S3R (ITS II). Numbers from 1 to 3 are replicas of DNA of each
 species. + is
 383 the positive control. – is the negative control. M is the 100 bp DNA Ladder (Cat. #
 62101,
 384 Promega).

Table 1 (on next page)

Primers used for amplification of *rbcL*, *matK*, ITS1 and ITS2

1 Table 1. Primers used for amplification of *rbcL*, *matK*, ITS1 and ITS2.

Primer pairs	Sequence	Size (bp)	Locus	Reference
rbcLA_F/ rbcLA_R	ATGTCACCACAAACAG AGACTAAAGC GTAAAATCAAGTCCAC CRCG	550	rbcL	Costion et al. 2011
matK_3F_KIM f/matK_1R_KIM R	CGTACAGTACTTTTGTG TTTACGAG ACCCAGTCCATCTGGA AATCTTGGTTC	850	matK	Costion et al., 2011
ITS 5a F/ ITS 4 R	CCTTATCATTTAGAGGA AGGAG TCCTCCGCTTATTGATA TGC	700	ITS1	Schultz et al. 2005
S2F/ S3R	ATGCGATACTTGGTGT GAAT GACGCTTCTCCAGACT ACAAT	400	ITS2	Schultz et al. 2005

2
3

Table 2 (on next page)

Samples and sequences submitted in the GenBank from the samples of *M. coriacea* barcoded

1 Table 2. Samples and sequences submitted in the GenBank from the samples of *M.*
coriacea

2 barcoded.

Barcode	Accession
<i>rbcL</i>	2198607
<i>matK</i>	2199742
ITS1	MK577640
ITS2	MK577643

3

Table 3(on next page)

Best model to describe the substitution pattern using Mega7

1 Table 3. Best model to describe the substitution pattern using Mega7.

Barcode	Best model
<i>rbcL</i>	JC
<i>matK</i>	T92
ITS1	T92+G
ITS2	T92+G

2 KG: Kimura 2-parameter; +G: Gamma distribution; T92: Tamura 3-parameter; GTR: General

3 Time Reversible. K2: Kimura 2-parameter. JC: Jukes-Cantor.

4

Figure 1 (on next page)

Macro morphological details of leaf from *M. coriacea*

A: retuse apex, B: whole edge, C: obtuse base, D, E and F: closed rib

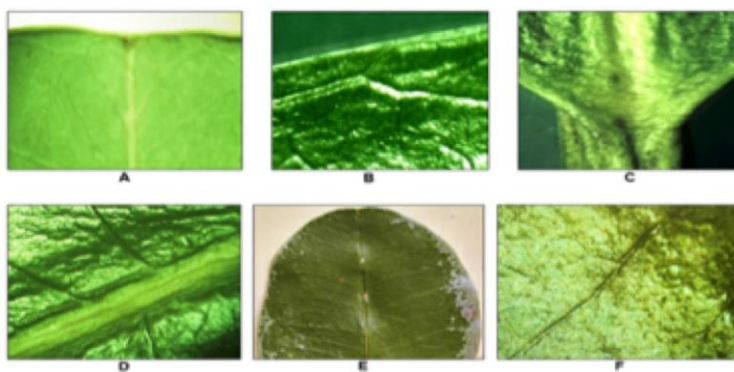


Figure 2(on next page)

Macro morphological details of crust from *M. coriacea*

A: external surface, **B:** internal surface



Figure 3(on next page)

Macro morphological characters of fruits and seeds from *M. coriacea*

a) green fruit, b) ripe fruit, c) seeds green fruits with peel, d) seeds ripe fruits with peel, e) endosperm green seeds, f) endosperm mature seeds

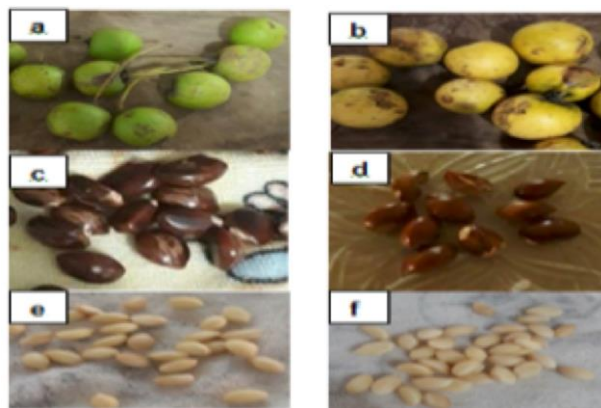


Figure 4

Microscopic characteristics of leaf from *M. coriacea*

Transversal section of the central nerve of the leaf (I): A: central nerve of the leaf, B and C: enlarged view of the central nerve, D: mesophylic, Cu: cuticle, Ep: epidermis, COC: calcium oxalate crystals, SP: spongy parenchyma, VS: vascular system, En: endodermis, AdE: adaxial epidermis, PP: palisade parenchyma, AbE: abaxial epidermis.

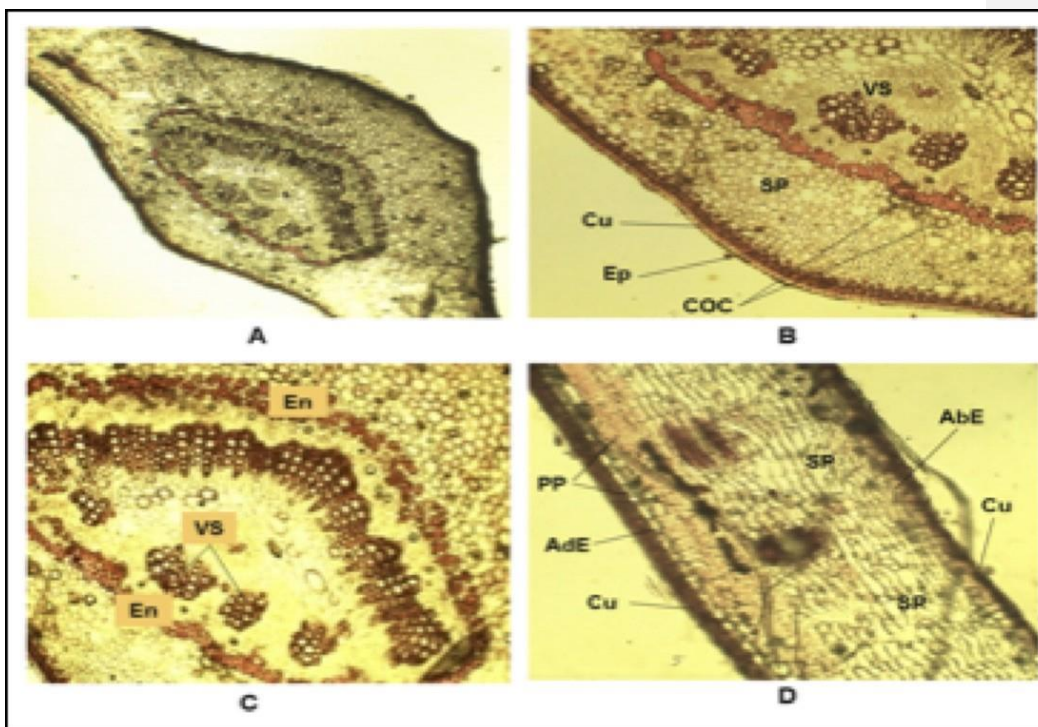


Figure 5(on next page)

Microscopic characteristics of leaf from *M. coriacea*

Diafanized of the leaf (II): E: adaxial epidermis, F and G: abaxial epidermis EpC: epidermal cells, S: stomata, EO: essential oils

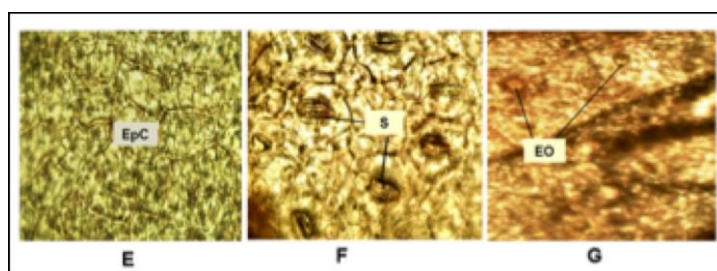


Figure 6

Powder drug characteristics of *M. coriacea*

VS: vascular system, F: fibers, S: starch, ST: suberous tissue, SF: septate fibers, COC: calcium oxalate crystal, SC: sclerides cells, MS: macrosclerides, OB: oilbag, SG: starch granules

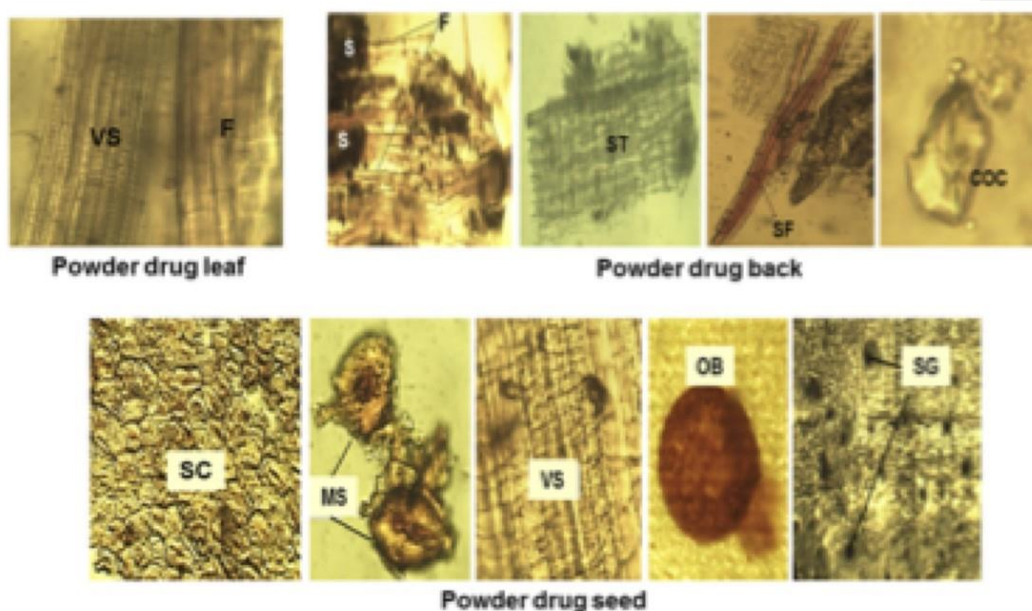


Figure 7 (on next page)

Gel electrophoresis of amplicons generated for the molecular barcodes with the genomic DNA of *M. coriacea*

A amplification of rbcLA_F/ rbcLA_R. **B** Amplification of matK_3F_KIM f/matK_1R_KIM R. **C** amplification of ITS 5a_F/ITS 4_R (ITS I). **D** amplification of S2f/S3R (ITS II). Numbers from 1 to 3 are replicas of DNA of each species. + is the positive control. – is the negative control. M is the 100 bp DNA Ladder (Cat. # 62101, Promega).

