Molecular identification using DNA barcoding and pharmacognostical evaluation of *Justicia beddomei* (C. B. Clarke) Bennet

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**ARTICLE HISTORY**
Received on: 10/10/2023  
Accepted on: 22/12/2023  
Available Online: XX

**Key words:**  
*Justicia beddomei* (C.B. Clarke) Bennet, trnH–psbA barcode, NCBI database, pharmacognostical studies, molecular identification, DNA sequence.

**ABSTRACT**
*Justicia beddomei* (C.B. Clarke) Bennet has been utilized for many years in the traditional medical system. This study aimed to facilitate the identification of *J. beddomei* (C. B. Clarke) Bennet using trnH–psbA DNA barcode region, NCBI database, and pharmacognostical features of the plant parts. Genomic DNA was extracted from plant leaf samples and polymerase chain reaction amplification and DNA sequence determination were performed. The contigs DNA sequence 319 bp was analyzed using the similarity basic search method BLASTn. The trnH–psbA barcode region of 319 bp contigs sequence showed 100% similarity with the standard sequence of *J. beddomei* with the Accession No. MK247216 from the NCBI database. Microscopical studies of the different parts of the plant facilitate the identification and differentiation of *J. beddomei* from its morphologically similar and confusing plant *Justicia adhatoda* L. *Justicia beddomei* can be easily identified by the flower and its inflorescence arrangement. The other identification features are the presence of cystoliths in the mesophyll region of the leaf, and the presence of colored content and crystals in the powder microscopic analysis. The outcome of the present detailed microscopic study of the stem, leaf, and flower of *J. beddomei* is of great value in the identification of the powdered samples and their adulterants.

**INTRODUCTION**
A large number of people in developing countries are approaching herbal species for their medicinal needs because of their safety over synthetic drugs. There is an equal range of international markets for herbal drugs and pharmaceutical products [1]. India is a rich source of herbal plants. Most human and animal diseases can be treated and prevented with the help of herbal plants [2,3]. Approximately 80% of the world’s population still rely on medications from herbal formulas [4]. The adulteration of herbal medicinal materials is becoming a global concern, coinciding with the growth of the herbal market. Therefore, more research should be supported to demonstrate the efficacy and security of natural herbal products [5]. Anatomical characteristics can be used to identify a small portion of a significant herbal product that is sold commercially as well as the actual plant source. The characteristics may be helpful in positively identifying plants [6].

DNA barcoding is a technique for identifying species depending on the analysis of short, uniform, and universal DNA region(s) (also known as “barcode sequence(s)”). This technique is widely used to identify a species by comparing the sequence of an unknown item to barcodes in a sequence database of recognized species [7,8]. Because of their physical resemblance to other plants in the same known species that are challenging to distinguish can be authenticated using DNA barcoding [9]. By comparing new species’ sequences to assembled reference libraries of barcode sequences, this technique enables the identification of new species. Many DNA sections, including matK, rbcL, trnH-PsbA, and ITS2, have been investigated for potential use as DNA barcodes [10].

The process of identifying, describing, and assessing the quality and purity of natural goods and plant-based
medicines is known as pharmacognostic evaluation. *Justicia* species are essential to many drug formulations used in Indian traditional medicine. “Vasa,” among other drug preparations, is a significant drug in Ayurveda. *Justicia adhatoda* and *Justicia beddomei* are reliable sources of “Vasa.” Particularly when dried, the morphological characteristics of *Justicia* species can be confusing. Intentionally or unintentionally, *J. adhatoda* and *J. beddomei* are replaced with species such as *J. carnea*, *J. betonica*, *J. gendarussa*, *J. santapau*, and *J. wynaadensis* when making the Ayurvedic medication “Vasa.” To evaluate the botanical identification, authenticity, purity, and quality of medicinal plant resources and their products, a variety of scientific procedures and approaches are applied [11–14].

In *Justicia* species, 56 different varieties are recorded in continental India [15]. Of these, 30 species are distributed in the Western Ghats regions and 26 species are present in Kerala [16]. *Justicia beddomei* (C.B. Clarke) Bennet [syn. *Adhatoda beddomei* C.B. Clarke] is a member of the Acanthaceae family [14]. Treatment of colds, coughs, asthma, bronchitis, and tuberculosis has long been practiced in India using tropical plants of the genus *Adhatoda* (Acanthaceae), usually referred to as “Vasa” [17]. The plant has many morphological traits with *J. adhatoda*, except for a few minor differences [14]. The main difference is in floral characteristics and similarities are in leaf and stem parts. The plant is an erect perennial shrub, 1.5–3 m tall; branches glabrous, terete to subtetragonal. This species is solely restricted to India’s Southern Western Ghats [17]. Recent literature has mentioned *J. beddomei* from Kerala as Chittadalodakam [14,17]. *Justicia beddomei* has historically been used to treat a number of illnesses, including fever, cough, leprosy, heart problems, blood disorders, hemorrhage, and tuberculosis. Some articles describe the Malayalam name as “Cheriya aadadalakam/Chittadalodakam for *A. beddomei* [18]” which is a variant of *Justicia adathoda* L and is not *J. beddomei* (C.B. Clarke) Bennet. The commonly found plant is *Justicia adathoda* L. *J. beddomei* is a different species with very similar macroscopic features to *J. adathoda* L. Therefore, the taxonomy experts found difficulty in authenticating this plant in the nonflowering season based on morphological characters [20]. The main identification features of the plant are the inflorescence and floral characteristics. The flowering season of the plant is between December and June. The current article proposes an alternate method to identify the plant in the nonflowering season by DNA barcoding technique and also updates the information regarding the main microscopic difference of various parts of *J. beddomei* with *J. adathoda* L [17].

Despite the fact that *J. beddomei* (C.B. Clarke) Bennet (Fig. 1) contains a lot of documented phytochemical information and is listed in Critically Endangered Category [18,20]. There are only a few molecular studies on the *Justicia* species using ITS and matK genes [21]. There is no proof of its molecular identification using trnH-PsbA DNA barcode, detailed pharmacognostic (macroscopic and microscopic) features of various parts of this plant (leaf, stem, and flower), and power microscopic features of the whole plant of this plant. This study will facilitate the identification of this plant and improve the identification of adulterants in further studies of this plant.

**Figure 1. Justicia beddomei** (C.B. Clarke) Bennet.

### MATERIALS AND METHODS

**Plant material collection and authentication**

The plant, *J. beddomei* (C.B. Clarke) Bennet was taken from the herb garden at the Arya Vaidya Sala in Kottakkal. The plant was identified by the Department of Forest Botany, Kerala Forest Research Institute (KFCRI), to be *J. beddomei* (C.B. Clarke) Bennet. The plant specimen’s herbarium is saved at Kerala Forest Research Institute (Herbarium), Peechi, Thrissur (Accession Number-19373).

**Chemicals**

All analytical grade chemicals used were commercially purchased from Chemind, Thrissur, Kerala.

**Sample preparation for DNA extraction**

Young, healthy leaves from the selected plant species were used to prepare the samples. After collection, the samples were produced using the sample identification number one. Leaves were professionally washed, dried, and stored at room temperature in sealed plastic packets [22]. Leaf samples were cut into tiny pieces after the midribs were removed. A mortar and pestle were used to grind 0.4 g of the sample into a powder using liquid nitrogen. The stored samples were later utilized to extract DNA in the molecular biology laboratory.

**DNA extraction**

DNA was isolated from a 0.4 g sample of leaf tissue using the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany) in accordance with the manufacturer’s instructions. Gel electrophoresis is used on a 0.8% agarose gel with 5 µl of sample standard DNA loaded (Sample M) per each lane, the quality of the isolated DNA was evaluated by a spectrophotometric approach using the Bio photometer plus (Eppendorf, Germany).

**Polymerase chain reaction (PCR) amplification and purification**

The PCR amplification was carried out using a 20 µl reaction mixture containing 5–20 ng of DNA: 1X PCR buffer (2
mM MgCl2), 200 µM each of dATP, dCTP, dGTP, and dTTP, 0.5 µM of each forward and reverse primer, 1U of Taq polymerase (Takara, Japan). The DNA amplification was carried out in a thermal cycler (Eppendorf, Germany). The list of primers and their nucleotide sequences are shown in Table 1. For 35 cycles of the PCR, the temperature was set at 92°C for 4 minutes, 94°C for 1 minute, 52°C for 1 minute, 64°C for 1 minute, and then 64°C for an additional 8 minutes. The 10 µl PCR reaction combination was used [23,24]. The tubes were taken out of the thermal cycler once the reaction was complete and kept at 4°C for storage. Gel-loading dye and PCR products were mixed, and the outcomes were separated using 1.8% agarose gel electrophoresis. The PCR products were purified after they had been separated on an agarose gel using the Machery-Nagel, Germany-based NucleoSpin® Gel, and PCR Clean-up kit. The purified PCR products were sequenced on an ABI 3730xl DNA sequencer at the AgriGenome laboratories facility in India for bidirectional sequencing (Sanger DNA sequencing) using M13Forward or M13Reverse primers, since all of our particular F/R primers were connected to M13F and M13R sequences, respectively.

Sequence editing, alignment, and analysis

The software, BioEdit sequence alignment editor version 7.2.6, was used to manually alter the unprocessed DNA sequences [25]. The CLUSTALW [26] algorithm in BioEdit was used to align the forward and reverse sequences, and the alignments were manually reviewed and corrected. The overlapping regions were joined together to obtain the contiguous DNA sequences [27]. The sequences were analyzed using the similarity-based method BLASTn available from the National Center for Biotechnology Information (www.ncbi.nlm.nih.gov). NCBI BLASTn to find homologies between the fragments and identify species [28].

Pharmacognostical evaluation

Organoleptic and macroscopic evaluation

It is referred to as the evaluation of plant material employing elements such as color, odor, taste, form, and texture. Various parts leaf, stem, and flower of the plant were considered for the macroscopical study [29].

Microscopic evaluation

Cross sections of live plant materials from the leaf, stem, and flower portions were cut, and each part separately under a microscope. Microscopic features of plant powder were analyzed [30,31]

Table 1. List of primers used for amplification/sequencing of trnH-psbA locus.

<table>
<thead>
<tr>
<th>DNA region</th>
<th>Primers name</th>
<th>Sequence (5′-3′)</th>
</tr>
</thead>
<tbody>
<tr>
<td>trnH-psbA</td>
<td>psbA3_f</td>
<td>GTTATGCATGAA CGTAATGCTC</td>
</tr>
<tr>
<td></td>
<td>trnHf_05</td>
<td>CGCGCATGGTGG ATTCACAATCC</td>
</tr>
<tr>
<td>Sequecing</td>
<td>M13F (-21)</td>
<td>TGTAAAACGACG GCCAGT</td>
</tr>
<tr>
<td>primers</td>
<td>M13R (-27)</td>
<td>CAGGAAACAGCT ATGACC</td>
</tr>
</tbody>
</table>

RESULT AND DISCUSSION

PCR amplification

Using the trnH-psbA region, the juvenile leaf of the J. beddomei genome was characterized. The trnH-psbA intergenic spacer region has been used in many DNA barcoding studies. This area of the plant was amplified by the primers psbA3_f and trnHf_05. Figure 2 shows that the nuclear trnH-psbA barcode sections of the sample were amplified using universal primers with 100% PCR performance after DNA extraction and nuclear trnH-psbA region PCR amplification in plant J. beddomei (C.B. Clarke) Bennet. A clear band of about 400 bp in length was visible in the sample (Fig. 2). All the primers amplified a specific target region of 400 bp.

Sample 1 amplified a ~400 bp product using the trnH-psbA primers. After sequencing the amplified products, the final DNA sequence contigs contain 319 bp. The obtained sequence was analyzed in the NCBI—BLASTn for species identification by comparing the nucleotide query sequence to a nucleotide sequence database. The top 10 results obtained were selected (Table 2). From the results, trnH-psbA barcode region of the selected plant was compared with the BLASTn and showed 100% similarity with the plant J. beddomei nucleotide NCBI database with Accession no: MK347214.1. The BLASTn result’s percentage of identity verified the taxonomic identity of the plant, indicating that the plant was successfully identified by the DNA barcoding method.

Pharmacognostical evaluation

Macroscopy

The leaves of the plant are light green in color, simple, opposite, and exstipulate with the short petiole (Fig. 3A). Leaf
shape is oblong-lanceolate. Leaves are entire and glabrous. Leaves measure 7.2–14.1 cm in length and 0.8–3.5 cm in breadth. The inflorescence is a short peduncled, condensed spike, 4.5–5 cm in length. Spike is axillary and stout. The leaf of the morphologically similar plant *J. adhatoda* is in dark green color and possesses a long petiole [14].

A small piece of stem is separated from a leafed twig (Fig. 3B). It measures 6.1 cm in length and 0.5 cm in width. The stem is green in color. It is unbranched, quadrangular, and thick. It is hard to touch, rough and woody texture. It has several brown patches. The odor of the plant is not characteristic and is bitter in taste.

The flower is the main identification part of the plant. Flowering season is from the month of December to June. Flowers are zygomorphic, bisexual, sessile, and 1.5 cm long (Fig. 3C and D) and the inflorescence, number of flowers, is 4. The flower of *J. adhatoda* plant is 3–3.5 cm long and the inflorescence contains a minimum of 10 flowers [14].

**Microscopy**

**Leaf**

*Justicia beddomei*’s transverse section is slightly depressed centrally, on the upper side and convex on the bottom side. Upper and lower surfaces have a single-layered epidermis which is square-to-rectangular, radially organized cells with a cuticle. A few simple, uniserate, long, straight or bent, thick-walled nonglandular covering trichomes were found. Sessile, four-celled glandular trichomes were also found. Hypodermis is collenchymatous, tightly packed followed by parenchymatous ground tissue. The midrib and laminar region contains calcium oxalate crystals. Vascular bundles are located at the center of the midrib composed of the xylem and phloem. Vascular bundles include 3–4 conjoint, collateral meristoles. Centrally located arc-shaped meristele is the largest among them. Smaller meristoles are located on both lateral sides (Fig. 4).

The laminar region is composed of the upper epidermis, mesophyll layer, and lower epidermis. The mesophyll layer is composed of palisade and spongy parenchyma (Fig. 5). Palisade parenchyma is thin-walled, columnar cells, found below the upper epidermis followed by circular to oval-celled spongy parenchyma found in the lower epidermis. Oil globules are found in the mesophyll region. Crystals of calcium oxalate and elongated warty cystoliths are also present in the mesophyll region. Stomata is diacytic and is found more in the lower epidermis than in the upper epidermis (Table 3).

The Transverse section of *J. adhatoda* leaf [32] is almost similar to *J. beddomei*. The main difference is in the cystolith region. The cystolith part extends from the epidermis to the lower cells of the palisade in *J. adhatoda*. Whereas, in *J. beddomei* cystolith part is in the mesophyll region. The stomatal index of *J. beddomei* is similar to *J. adhatoda*.

**Stem**

The transverse section of *J. beddomei* stems shows cork, followed by a parenchymatous cortex. Crystals of calcium oxalate are found in the collenchymatous hypodermis. A band of stone cells, mostly rectangular in shape with distinct pits are also seen in the cortical region. The xylem is composed of lignified vessels, fibers, parenchyma, and rays. Phloem is

**Table 2. The top 10 BLASTn results from the NCBI database.**

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Scientific name</th>
<th>Max score</th>
<th>Total score</th>
<th>Query cover</th>
<th>E value</th>
<th>% identity</th>
<th>Acc. Length</th>
<th>Accession</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><em>Justicia beddomei</em></td>
<td>590</td>
<td>590</td>
<td>100%</td>
<td>3.00E⁻¹⁴⁴</td>
<td>100</td>
<td>319</td>
<td>MK347214.1</td>
</tr>
<tr>
<td>2</td>
<td><em>Justicia adhatoda</em></td>
<td>366</td>
<td>366</td>
<td>100%</td>
<td>6.00E⁻⁹⁷</td>
<td>87.01</td>
<td>375</td>
<td>MG947188.1</td>
</tr>
<tr>
<td>3</td>
<td><em>Justicia adhatoda</em></td>
<td>366</td>
<td>366</td>
<td>100%</td>
<td>6.00E⁻⁹⁷</td>
<td>87.01</td>
<td>378</td>
<td>MG947088.1</td>
</tr>
<tr>
<td>4</td>
<td><em>Justicia adhatoda</em></td>
<td>366</td>
<td>366</td>
<td>100%</td>
<td>6.00E⁻⁹⁷</td>
<td>87.01</td>
<td>354</td>
<td>MK347215.1</td>
</tr>
<tr>
<td>5</td>
<td><em>Justicia adhatoda</em></td>
<td>361</td>
<td>361</td>
<td>100%</td>
<td>3.00E⁻⁹⁵</td>
<td>86.72</td>
<td>395</td>
<td>KC420641.1</td>
</tr>
<tr>
<td>6</td>
<td><em>Justicia adhatoda</em></td>
<td>344</td>
<td>344</td>
<td>100%</td>
<td>3.00E⁻⁹⁰</td>
<td>85.88</td>
<td>FL*</td>
<td>NC047476.1</td>
</tr>
<tr>
<td>7</td>
<td><em>Justicia adhatoda</em></td>
<td>344</td>
<td>344</td>
<td>100%</td>
<td>3.00E⁻⁹⁰</td>
<td>85.88</td>
<td>FL*</td>
<td>MN711720.1</td>
</tr>
<tr>
<td>8</td>
<td><em>Justicia adhatoda</em></td>
<td>344</td>
<td>344</td>
<td>100%</td>
<td>3.00E⁻⁹⁰</td>
<td>85.88</td>
<td>382</td>
<td>MG947151.1</td>
</tr>
<tr>
<td>9</td>
<td><em>Justicia adhatoda</em></td>
<td>303</td>
<td>303</td>
<td>89%</td>
<td>4.00E⁻⁷⁸</td>
<td>85.62</td>
<td>341</td>
<td>MG947201.1</td>
</tr>
<tr>
<td>10</td>
<td><em>Justicia adhatoda</em></td>
<td>303</td>
<td>303</td>
<td>61%</td>
<td>4.00E⁻⁷⁸</td>
<td>95.51</td>
<td>361</td>
<td>KR108262.1</td>
</tr>
</tbody>
</table>

*FL-Chloroplast full length sequence.*
towards the base. Anther is 2-celled and cells are superposed. The ovary is superior, light green, glabrous, bicarpellate. Style is filiform, pubescent and Stigma is minutely bifid, dull-white, sparsely pubescent (Fig. 8).

Microscopy transverse section

The transverse section (T.S.) of gynoecium is almost oval in outline. It shows a bicarpellary ovary with two locules. Each locule contains one ovule in each chamber. The ovary is hypogynous (Fig. 9). The main difference between *J. adhatoda* and *J. beddomei* is in the T.S of gynoecium, i.e., the presence of phloem elements. Medullary rays are also found in the vascular region. Pith is composed of pitted parenchyma. A number of raphids are seen in the pith region (Figs. 6 and 7).

Table 3. Stomatal index of *J. beddomei* leaf.

<table>
<thead>
<tr>
<th>Sl. No</th>
<th>Sample</th>
<th>Stomatal index (I) %</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Upper epidermis</td>
<td>2.22 ± 0.021</td>
</tr>
<tr>
<td>2</td>
<td>Lower epidermis</td>
<td>11.76 ± 0.016</td>
</tr>
</tbody>
</table>

Longitudinal section of flower

The longitudinal section of *J. beddomei* flower shows corolla, androecium, and gynoecium. Corolla is white, and imbricate. Corolla tube is pubescent inside, 2-lipped, the lower lip is 3-lobed, striations absent on the lip and the upper lip is bifid. There are two stamens, basifixed and light green. Filaments are dull white, and glabrous except for hirsute

![Figure 4. Transverse section of midrib of *J. beddomei*.](image)

![Figure 5. Transverse section of leaf lamina of *J. beddomei*.](image)

![Figure 6. Enlarged portion of *J. beddomei* stem.](image)

![Figure 7. Some characters of *J. beddomei* stem.](image)
of abundant unicellular to multicellular trichomes in the outer epidermis of \textit{J. adhatoda} \[31\] which is absent in \textit{J. beddomei}. The major difference in macroscopy and microscopic features of \textit{J. beddomei} and \textit{J. adhatoda} is illustrated in a tabular form (Table 4).

**Powder microscopy**

Powder microscopy of the whole plant of \textit{J. beddomei} powder shows spiral vessels, vessels with reticulated thickening, pitted vessels, a fragment of lignified and nonlignified fiber, a large number of diacytic stomata, and few crystals. Uniseriate, covering, straight trichomes and four-celled glandular trichomes are also found along with fragments of the mesophyll layer.

**Table 4. Difference between macroscopy and microscopic features of \textit{J. beddomei} and \textit{J. adhatoda}.**

<table>
<thead>
<tr>
<th></th>
<th>\textit{Justicia beddomei}</th>
<th>\textit{Justicia adhatoda} [16,32,34]</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Macroscopy</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leaf: Light green</td>
<td>Leaf: Dark green</td>
<td></td>
</tr>
<tr>
<td>Leaf: Short petiole</td>
<td>Leaf: Long petiole</td>
<td></td>
</tr>
<tr>
<td>Flower: 1.5 cm long</td>
<td>Flower: 3–3.5 cm long</td>
<td></td>
</tr>
<tr>
<td>Ovary: Glabrous</td>
<td>Ovary: Pubescent</td>
<td></td>
</tr>
<tr>
<td>Inflorescence: Number of flowers</td>
<td>4–6</td>
<td>Inflorescence: Number of flowers</td>
</tr>
<tr>
<td><strong>Microscopy</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leaf: Lower epidermis contain nonglandular trichomes</td>
<td>Leaf: Lower epidermis contain glandular trichomes</td>
<td></td>
</tr>
<tr>
<td>Leaf: Elongated warty cystolith in mesophyll region</td>
<td>Leaf: Cystolith extend from epidermis to lower cells of palisade.</td>
<td></td>
</tr>
<tr>
<td>Gynoecium: Absence of abundant unicellular to multicellular trichomes</td>
<td>Gynoecium: presence of abundant unicellular to multicellular trichomes</td>
<td></td>
</tr>
</tbody>
</table>

Colored contents, fragments of pitted parenchyma, sclerids, and stone cells are also found (Fig. 10).

Powder microscopy of \textit{J. beddomei} and \textit{J. adhatoda} showed prismatic calcium oxalate crystals \[17\]. Different epidermal trichomes exist in both plants \[34\]. In \textit{J. beddomei}, only glandular trichomes are observed, whereas, in \textit{J. adathoda}, both glandular and nonglandular trichomes are visible. \textit{Justicia beddomei} is the sole species with the colored content.

**CONCLUSION**

This study revealed the identification concerns of the plant \textit{J. beddomei} (C.B. Clarke) Bennet with the morphologically similar plant \textit{J. Adathoda}. The sequence analysis and NCBI Gen-Bank BLASTn database results after the amplification of the \textit{trnH–psbA} barcode region of the plant facilitated the identification of this species. The distinguished morphological and microscopical features of \textit{J. beddomei} (C.B. Clarke) Bennet are the four number of flowers in inflorescence,
1.5 cm flower length, and presence of cystolith in the mesophyll region of leaf; absence of trichomes in gynoecium, presence of glandular trichomes only in power microscopy, respectively. These features will help to find the adulteration in the selected plant studies and ensure the purity of the selected drug. The powder microscopic analysis helps to minimize the adulteration of drugs in powdered form. This article will be helpful for the preparation of J. beddomei (C.B. Clarke) Bennet monograph and pharmacopeial standards. The distinguished features in microscopy of leaf, stem, and flower gave relevant information for the identification of the plant from J. Adathoda L. According to our knowledge, this is the first thorough analysis of molecular identification using trnH-psbA DNA barcoding and thorough microscopic, macroscopic, and powder microscopic features of J. beddomei (C.B. Clarke) Bennet parts.

ACKNOWLEDGMENT

The authors would like to thank CMPR, Arya Vaidya Sala, Kottakkal, Kerala, India, for providing the necessary facilities to carry out the DNA barcoding analysis.

AUTHOR CONTRIBUTIONS

SP and NR conceptualized and designed the research work, NR significantly contributed to the practical part and initial drafting of the manuscript; SP revised the article examining it carefully for significant intellectual content; consenting to submit to the current journal; SP gave the version’s final drafting of the manuscript; SP gave the version’s final drafting of the manuscript; SP revised the article examining it carefully for significant intellectual content; consenting to submit to the current journal; SP gave the version’s final approval. All the authors are eligible to be authors as per the International Committee of Medical Journal Editors (ICMJE) requirements/guidelines

FINANCIAL SUPPORT

There is no funding to report.

CONFLICT OF INTEREST

The authors declare no conflicts of interest.

ETHICAL APPROVAL

This study does not involve experiments on animals or human subjects.

DATA AVAILABILITY

All data generated and analyzed are included in this research article.

PUBLISHER’S NOTE

This journal remains neutral with regard to jurisdictional claims in published institutional affiliation.

REFERENCES


How to cite this article: