Chemical Composition and Biological Activities of Essential Oils from Fresh *Vismia guianensis* (Aubl.) Choisy and *Vismia cayennensis* (Jacq.) Pers. Leaves

Composição química e atividades biológicas dos óleos essenciais das folhas de *Vismia guianensis* (Aubl.) Choisy e *Vismia cayennensis* (Jacq.) Pers.

Composición química y actividades biológicas de los aceites esenciales de hojas de *Vismia guianensis* (Aubl.) Choisy y *Vismia cayennensis* (Jacq.) Pers.

Received: 06/21/2021 | Reviewed: 06/29/2021 | Accept: 06/30/2021 | Published: 14/07/2021

Antonia Tavares Barbosa
ORCID: https://orcid.org/0000-0002-8441-2699
Universidade Federal do Amazonas, Brazil
E-mail: antoniatavares92@hotmail.com

Vitor Hugo Neves da Silva
ORCID: https://orcid.org/0000-0002-1184-2241
Universidade Federal do Amazonas, Brazil
E-mail: hugor_rs@hotmail.com

Bruna Yuka Koide da Silva
ORCID: https://orcid.org/0000-0002-5228-7520
Universidade Federal do Amazonas, Brazil
E-mail: brunaykoide@gmail.com

Aniele da Silva Neves Lopes
ORCID: https://orcid.org/0000-0002-3830-9631
Universidade Federal do Amazonas, Brazil
E-mail: aniele.neves16@gmail.com

Isabel Reis Guesdon
ORCID: https://orcid.org/0000-0002-1372-250X
Universidade Federal do Amazonas, Brazil
E-mail: isabelbio@gmail.com

Paulo José Sousa Maia
ORCID: https://orcid.org/0000-0003-3101-3712
Universidade Federal do Rio de Janeiro-Macae, Brazil
E-mail: pmldcb@gmail.com

Maxwell Adriano Abegg
ORCID: https://orcid.org/0000-0002-0328-1122
Universidade Federal do Amazonas, Brazil
E-mail: maxabegg@gmail.com

Geone M. Corrêa
ORCID: https://orcid.org/0000-0002-9458-8305
Universidade Federal do Amazonas, Brazil
E-mail: geonemaia@ufam.edu.br

Dominique Fernandes de Moura do Carmo
ORCID: https://orcid.org/0000-0002-8835-1619
Universidade Federal do Amazonas, Brazil
E-mail: dominiquefmc@ufam.edu.br

Abstract

The *Vismia* Vand. genus encompasses many species indigenous to the Amazon rain forest where they are popularly known as “Lacre” bark and leaves are widely employed by locals to treat dermatophytoses. The aim of this study was to investigate the chemical composition of essential oils (EOs) extracted from the aerial parts of the species *Vismia guianensis* (Aubl.) Choisy and *Vismia cayennensis* (Jacq.) Pers. and to assess their antimicrobial activity against the bacteria *Staphylococcus aureus* Rosenbach 1884 and *Escherichia coli* (Migula 1895) Castellani and Chalmers 1919 as well as the fungi *Candida albicans* (C.P. Robin) Berkhout 1923 and *Candida parapsilosis* (Ashford) Langeron & Talice 1932. The analysis of the chemical composition of the essential oil extracted from *V. guianensis* leaves (EOVg) indicated 46 components, of which three sesquiterpenes predominated, namely: (E)-caryophyllene (10.40%), α-copaene (29.45%), and (E)-nerolidol (24.06%). As to the essential oil from *V. cayennensis* leaves (EOVc), 61 components were identified, of which two oxygenated sesquiterpenes stood out as the main components, namely, germacrone (25.42%) and curzerene (25.29%). EOVg exhibited Minimum Inhibitory Concentration (MIC) of 1.56 μg/mL against the yeast *C. parapsilosis* whereas EOVc was active against the bacteria *E. coli* and *S. aureus* as well as the yeast *C. parapsilosis*. The results obtained in this study strongly recommend further research on the essential oils.
in question with a view to isolating and identifying the components responsible for their observed antimicrobial activities.

**Keywords:** Essential oils; Chemical composition; Hypericaceae.

**Resumen**

El género *Vismia* Vand. tiene especies distribuidas en la Región Amazónica, donde se les conoce popularmente como “Lacre” y su corteza y hojas se utilizan en el tratamiento de dermatofitosis. Este trabajo tuvo como objetivo estudiar la composición química de los aceites esenciales de las partes aéreas de las especies *Vismia guianensis* (Aubl.) Choisy y *Vismia cayennensis* (Jacq.) Pers. y evaluar la actividad antimicrobiana frente a las bacterias *Staphylococcus aureus* Rosenbach 1884 y *Escherichia coli* (Migula 1895) Castellani and Chalmers 1919 y los hongos *Candida albicans* (C.P. Robin) Berkhout 1923 y *Candida parapsilosis* (Ashford) Langeron & Talice 1932. Sobre la composición química de aceite esencial (AE), se identificaron 46 constituyentes en el aceite esencial de la especie *V. guianensis* (AEVg), de los cuales tres fueron identificados como mayores sesquiterpenos: trans-cariofileno (10,40%), α-copaeno (29,45%) y trans-nerolidol (24,06%). En la muestra de aceite esencial de la especie *V. cayennensis* (AEVc) se identificaron 61 constituyentes, de los cuales se destacaron dos sesquiterpenos oxigenados como constituyentes mayoritarios, germacrón (25,42%) y cariofileno (25,29%). O AEVg presentó Concentración Inhibitoria Mínima (CIM) de 1,56 µg/mL contra cepas fúngicas de *C. parapsilosis* y AEVc fue activo contra bacterias *E. coli* y *S. aureus* para la levadura *C. parapsilosis*. Los resultados obtenidos sugieren el aislamiento e identificación de componentes responsables las actividades observadas.

**Palabras clave:** Aceite esencial; Composición química; Hypericaceae.

1. **Introduction**

Essential oils from plants have been widely used in medicine, agriculture, perfumery, and cosmetics (Paolini, et al., 2010; Thuy, et al., 2021; Maia, et al., 2019; Křúmal, et al., 2015; Razavi, et al., 2021; Saikia, et al., 2020; Stojanović-Radić, et al., 2020). Several research groups are currently conducting studies on the chemical composition and biological potential of essential oils extracted from numerous plants, such as *Vismia* Vand. spp. (Simões et al., 2007). *Vismia* is an extensive genus of the Hypericaceae family, consisting of small trees inhabiting the tropical and subtropical regions of Central and South America (Hussain et al., 2012).

Notwithstanding the extensiveness of this genus, the literature provides scientific studies on essential oils from three *Vismia* species alone. Rojas et al. (2011) studied essential oils obtained from *V. guianensis* (Aubl.) Choisy fruit, *V. baccifera* Planch. & Triana fruit, and *V. macrophylla* Kunth leaves and reported antimicrobial activity for *V. baccifera* var. *dealbata* (Kunth) Ewan against the microorganisms *Enterococcus faecalis* (Andrewes and Horder 1906) Schleifer and Kilpper-Bälz 1984, *Staphylococcus aureus* Rosenbach 1884, and *Bacillus cereus* Frankland and Frankland 1887, with MIC values ranging from 9 to 37 µg/mL. Another study by Rojas et al. (2010) on the essential oil (EO) extracted from *V. baccifera* fruit pointed to broad-spectrum bactericide activity against *Staphylococcus aureus*, *Enterococcus faecalis*, *Escherichia coli* (Migula 1895) Castellani and Chalmers 1919, *Pseudomonas aeruginosa* (Schroeter 1872) Migula 1900, and *Klebsiella pneumoniae* (Schroeter ...
1886) Trevisan 1887. By means of CG-EM analysis, the researchers identified 27 chemical components in \textit{V. baccifera} EO, chiefly trans-cadin-1,4-diene, cis-cadin-1,4-diene, and \( \beta \)-caryophyllene.

Additionally, Rojas et al. (2011) described the fungicide potential of \textit{Vismia} essential oils, which supports the scientific community’s interest in identifying the chemical components and assessing the pharmacological actions of this genus. In this vein, this study aimed at investigating the chemical profile and antimicrobial potential of essential oils extracted from \textit{V. guianensis} and \textit{V. cayennensis} (Jacq.) Pers. leaves against the microorganisms \textit{E. coli}, \textit{S. aureus}, \textit{Candida albicans} (C.P. Robin) Berkhout 1923 and \textit{Candida parapsilosis} (Ashford) Langeron e Talice 1932.

\section{2. Materials and method}

\subsection*{2.1 Collection of plant samples}

Dry leaves from the aerial parts of the species under investigation were collected at several sites in Itacoatiara, Brazil: \textit{Vismia cayennensis} (03°05’36.9”S, 058°27’44.7”W Gr longitude, and ±5 m altitude) and \textit{V. guianensis} (03°05’39.8”S, 058°27’43.8”W Gr longitude, ±4 m altitude). Then, the samples were identified and deposited at the herbarium collection of the Universidade do Estado do Amazonas, at the Centro de Estudos Superiores de Itacoatiara–CESIT (Voucher number 0055) and at the Instituto Nacional de Pesquisas da Amazônia-INPA (Voucher number 18448).

\subsection*{2.1.1 Extraction of essential oils from \textit{Vismia guianensis} and \textit{V. cayennensis} leaves}

The essential oils from \textit{V. guianensis} and \textit{V. cayennensis} leaves were obtained by hydrodistillation of fresh material in a modified Clevenger-type apparatus coupled to a round bottom flask with distilled water (5 L) for approximately 4 hours. Next, the EO samples were centrifuged (3500 rpm) for 10 minutes for water/oil separation, which was done with a graduated micropipette. Extractions were performed in triplicate, with 800 g of fresh leaves of both \textit{Vismia} species in each flask. Then, the EO samples were stored in Eppendorf tubes, sealed, and kept at -4 °C until analysis and testing. The samples of \textit{V. guianensis} and \textit{V. cayennensis} EOs were coded as EOVg and EOVc, respectively.

\subsection*{2.1.2 Gas chromatography coupled to mass spectrometry (GC/MS)}

The obtained essential oils underwent analysis in a SHIMADZU gas chromatographer coupled to a SHIMADZU QP2010 mass spectrometer (GC/MS). For component chromatography, a 30 m \( \times \) 0.25 mm DB-5MS column with 0.25 \( \mu \)m inner film thickness was employed. The chemical components were identified by interpreting their respective mass spectra, calculating their Kovats Indexes (KIs), and matching them up to data found in the literature. The calculated KIs were then compared to those tabulated for isolated compounds by Adams (2007).

\subsection*{2.2 Biological assays}

\subsection*{2.2.1 Antimicrobial assay}

The antimicrobial activity test was carried out at the Microbiology Laboratory of the Exact Sciences and Technology Institute (ICET/UFAM), using the microplate dilution test to analyze the antibacterial and antifungal potential of EOVg and EOVc.

\subsection*{2.2.2 Microorganisms}

The EO samples underwent antimicrobial susceptibility tests \textit{in vitro} according to the protocol described in the literature (Vaz et al., 2009), using a panel with ATCC strains (American Type Culture Collection, USA). The antibacterial
activity of the essential oils was assessed against the Gram-positive bacterium *Staphylococcus aureus* (ATCC 25923), Gram-negative bacterium *Escherichia coli* (ATCC 25922), and fungi *Candida albicans* (ATCC 10231) and *Candida parapsilosis* (ATCC 22019).

### 2.2.3 Preparation and standardization of microbial inoculums

The bacterial and fungal strains were grown in Mueller Hinton broth for 24 h at 37 °C and standardized by adding sterile PBS (pH 7.2) until obtaining turbidity equal to that of the suspension in the 0.5 tube on the McFarland scale (approximately 1.0 × 10⁸ CFU/mL). Then, a spectrophotometric reading was performed at 620 nm to confirm the microorganism concentration. Subsequently, small amounts of bacterial and fungal strains were removed, with the aid of a sterile loop, and added to 5 mL of sterile LB broth and 5 mL of YPD broth for the bacteria and fungi in question, respectively. The microorganism concentration was confirmed by spectrophotometer reading.

### 2.2.4. Preparation of EO samples

Firstly, a 10% dimethyl-sulfoxide (DMSO) solution was prepared by diluting DMSO (100 µL) in sterile distilled water (900 µL). Then the EOVg and EOVc samples were diluted in 10% DMSO by solubilizing the samples (10 mg) in the previously prepared solvent (100 µL). Through this procedure, stock solutions of each sample were prepared to a concentration of 100 µg/mL. The assays were performed on five 96-well ELISA microplates, which were divided as follows:

1. The wells used validating the method and measurement of results were identified as “positive and negative controls”: the positive control comprised the culture medium, bacterial or fungal suspension and the reference antimicrobial standard whereas the negative control consisted of the culture medium and 10% DMSO;
2. Wells identified as “blank” contained the culture medium and the essential oil of each sample in order to eliminate the turbidity caused by its color when evaluating the results; and
3. Wells identified as “assay” contained the culture medium, a mixture of essential oil with DMSO, and the bacterial or fungal suspension.

### 2.3 Determination of the Minimum Inhibitory Concentration (MIC)

#### 2.3.1 Microdilution for bacteria

MIC was performed in triplicate at 1:2 concentration. The bacteria under investigation were *E. coli* and *S. aureus*. Chloramphenicol and 10% DMSO were employed as reference standard (positive control) and negative control, respectively. The 96 microplate wells were filled with the LB broth (100 µL). Then, in the first well, 100 µL of the EO stock solution was added prepared initially in concentration at 100 µg/mL. After a serial dilution was conducted in the seven consecutive wells, removing 100 µL from the highest concentration well, resulting in a solution of up to 0.39 µg/mL. The assay was performed in triplicate for each concentration. Likewise, the LB broth (50 µL) plus Chloramphenicol (50 µL) were added to the positive control whereas the LB broth (50 µL) plus 10% DMSO (50 µL) were added to the negative control, in triplicate. Finally, the microorganism suspensions (10 µL) were added to every well and incubated for 24 hours at 37 °C.

#### 2.3.2 Microdilution for fungi

MIC was performed in triplicate for *C. albicans* and *C. parapsilosis* with methanol (positive control) and 10% DMSO (negative control) as reference standard. The microplate wells were filled with the YPD broth (100 µL). Then, in the first well, 100 µL of the EO stock solution was added prepared initially in concentration at 100 µg/mL. After a serial dilution was conducted in the seven consecutive wells, removing 100 µL from the highest concentration well, resulting in a solution of up to
0.39 µg/mL. The YPD broth (50 µL) plus methanol (50 µL) were added to the positive control whereas the YPD broth (50 µL) plus 10% DMSO (50 µL) were added to the negative control, in triplicate. Then, the microorganism suspensions (10 µL) were added to the wells and incubated for 48 h at 37 °C. After incubation, a visual reading and a reading with Resazurin dye (100 µg/mL) of the microbial growth were performed.

3. Results and discussion
3.1 Yields

Table 1 shows the variation in yields of EOVg and EOVc. EOVg yield was higher in September/2019 (0.04%), a period of intense drought, and lower yield in November/2019 (0.03%), the rainy season in the region. EOVc exhibited a yield of 0.54% in the dry season, a satisfactory value as compared to that obtained for EOVg during the same period. The difference between the yield values may be attributed to several factors, e.g., temperature, rainfall, place and time of sample collection, which can have critical effects on both the quantity and quality of essential oils.

| Plant sample       | Year/month | Temperature | Sample code | Plant mass (g) | Yield   |
|--------------------|------------|-------------|-------------|----------------|---------|
| V. guianensis      | September 2019 | 32 °C       | EOVg        | 800 g          | 0.04%   |
|                    | November 2019  | 30 °C       |             | 800 g          | 0.03%   |
| V. cayennensis     | September 2019 | 35 °C       | EOVc        | 800 g          | 0.54%   |
|                    | November 2019  | 30 °C       |             | 800 g          | 0.24%   |

* EOVg: Essential oil from Vismia guianensis; EOVc: Essential oil from V. cayennensis. Source: Authors.

The lower yields of the essential oils during the rainy season may be due to lixiviation, i.e., continuous rain may result in loss of hydro-soluble substances in leaves and roots. This may apply to plants that produce alkaloids, glycosides, and even volatile oils (Evans, 1996; Walteman & Mole 1994).

3.1.1 Chemical composition of essential oils

The analysis of the chemical composition of OEVg revealed 46 components, of which sesquiterpenes were the most common ones (63.91%): (E)-caryophyllene (10.40%), α-copaene (29.45%), and (E)-nerolidol (24.06%). OEVc presented 61 chemical components, in which a high proportion (50.71%) of oxygenated sesquiterpenes was found: germacrone (25.42%) and curzerene (25.29%). The chemical profile of both essential oils showed a high percentage of sesquiterpenes, mostly hydrocarbon sesquiterpenes. For example, V. guianensis has α-humulene (2.84%), β-selinene (0.79%), α-guaiene (0.16%), cyperene (0.20%), α-selinene (1.14%), α-murolene (0.23%), and δ-cadinene (2.27%) as well as two oxygenated sesquiterpenes, namely, caryophyllene oxide (0.88%) and viridiflorol (0.41%). In addition to this class of metabolites, hydrocarbon monoterpens, such as α-pinene (0.09%) and β-pinene (0.08%), and other components were identified, albeit in lesser quantities (Table 2).
were identified viridiflorol (0.45%), spatulenol (0.86%), sesquiterpene (0.04%), myrcene (0.24%), limonene (0.10%), and terpinolene (0.02%). The analysis of the monoterpenes, sesquiterpene, and carophyllene oxide showed the presence of α-pinene (2.15%), e.g., carophyllene (2.15%), and β-selinene (0.42%), as well as oxygenated sesquiterpenes, e.g., viridiflorol (0.45%), spatulenol (0.86%), and carophyllene oxide (0.97%). Likewise, small percentages of other components were identified, as shown in Table 3.

The analysis of the EOVC indicated the presence of hydrocarbon monoterpenes, such as α-pinene (0.08%), sabinene (0.04%), myrcene (0.24%), limonene (0.10%), and terpinolene (0.02%). Higher percentages were found for hydrocarbon sesquiterpenes, e.g., as (E)-carophyllene (2.15%), and β-selinene (0.42%), as well as oxygenated sesquiterpenes, e.g., viridiflorol (0.45%), spatulenol (0.86%), and carophyllene oxide (0.97%). Likewise, small percentages of the other components were identified, as shown in Table 3.
The main components of Vismia cayennensis EO were identified as α-pinene (36.6%), cadina-1,4-diene (36.6%), cadina-1,4-diene (18.8%), and β-caryophyllene (11.9%). They also identified β-caryophyllene (20.1%), germacrone D (11.6%), and β-elemene (7.0%) as the main components in a sample of essential oil extracted from V. macrophylla leaves. Mono- and sesquiterpenes have been identified as biologically active in the literature. Lima et al. (2005), investigating the action of the monoterpenes α-pinene and β-pinene, found satisfactory activity against C. albicans, C. tropicalis (Castell.) Berkhout, C. parapsilosis, C. stellatoidea (C.P. 71.21

| Substance | Class | Area % | RT | KI_{cal} | KI_{lab} |
|-----------|-------|--------|----|----------|----------|
| α-pinene  | HM    | 0.08   | 4.93 | 931      | 932      |
| Sabinene  | HM    | 0.04   | 6.067| 976      | 969      |
| Myrcene   | HM    | 0.24   | 6.330| 987      | 988      |
| Limonene  | HM    | 0.10   | 7.581| 1038     | 1024     |
| β-fandandrene | HM | 0.05   | 7.642| 1040     | 1025     |
| (Z)-β -ocimene | HM | 0.21   | 8.124| 1060     | 1044     |
| terpinolene | OM   | 0.02   | 9.516| 1116     | 1086     |
| δ-elemene | HS    | 0.33   | 19.698| 1329     | 1335     |
| (E)-caryophyllene | HS | 2.15   | 23.159| 1414     | 1417     |
| α-humulene | HS    | 0.22   | 24.626| 1450     | 1452     |
| germacrene-D | HS | 0.76   | 25.670| 1476     | 1484     |
| β-selinene | HS    | 0.42   | 25.980| 1483     | 1489     |
| curzerene | OS    | 25.29  | 26.290| 1491     | 1499     |
| muuroloene | HS    | 0.20   | 26.434| 1494     | 1500     |
| γ-cadinene | OS    | 0.42   | 27.192| 1513     | 1513     |
| spathulenol | OS    | 0.85   | 29.455| 1568     | 1577     |
| germacrone | OS    | 25.42  | 32.742| 1644     | 1693     |
| caryophyllene oxide | OS | 0.97   | 29.622| 1572     | 1482     |
| viridifloral | OS   | 2.44   | 29.799| 1577     | 1592     |
| cubeban-11-ol | -    | 0.96   | 30.221| 1587     | 1595     |
| Rosafiol | OS    | 0.45   | 31.259| 1612     | 1600     |
| δ-cadinene | HS    | 0.54   | 31.968| 1630     | 1522     |
| cadin-4-eno-7-ol <cis> | HS | 0.98   | 32.049| 1632     | 1635     |
| α-muurolool | HS    | 0.30   | 32147| 1634     | 1500     |
| hexadienoic acid | - | 0.88   | 32.365| 1631     | NI       |
| intermedeol | OM    | 5.28   | 32.560| 1642     | 1665     |
| naphthalenol | -    | 0.40   | 32.930| 1649     | NI       |
| 1,2,4,5 tetramethyl-[2,4] heptane | -   | 0.48   | 33.406| 1666     | NI       |
| (E)- γ --bisabolene | HS   | 0.73   | 42.589| 1890     | 1529     |

| Total     | 71.21 |

RT: Retention Time; KI_{cal}: Calculated Kovats Index; KI_{lab}: Tabulated Kovats Index; NI: Not Identified; HS: Hydrocarbon Sesquiterpene; OS: Oxygenated Sesquiterpene; OM: Oxygenated Monoterpene; HM: Hydrocarbon Monoterpene. Source: Adams (2017).
Jones & D.S. Martin) Langeron & Guerra, C. kruzei (Castell.) Berkhout, and Cryptococcus neoformans (San Felice) Vuill. In a previous study carried out by Peñuelas et al. (2005), monoterpenes were shown to have thermotolerance, photoprotection, and antioxidant properties due their ability to capture oxygen radicals produced during photosynthesis. On the other hand, sesquiterpenes have been highlighted in several essential oils due to their strong odor and anti-inflammatory and antifungal power (Zheng et al., 1992).

3.2 Biological activity

3.2.1 Microdilution assay for fungi and bacteria

The literature indicates remarkable antimicrobial activity in essential oils from Vismia spp. According to Pérez and colleagues (2011) and Montanari and colleagues (2011), essential oils extracted from V. macrophylla fruit are active against Gram-positive (S. aureus and E. faecalis) and Gram-negative (E. coli) bacteria, with MIC values ranging from 150 µL/mL to 740 µL/mL. Those studies also observed antimicrobial activity in the essential oil extracted from leaves of the same species, effective against the Gram-positive bacteria S. aureus (100 µL/mL) and E. faecalis (500 µL/mL) as well as against the fungi C. albicans and C. kruzei (600 µL/mL each). Screening results for antifungal and antibacterial activities are depicted in table 4.

| Table 4. Antibacterial and antifungal activities (MIC values in µg/mL) of OEVg and OEVc. |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Species         | Essential oils  | Bacterial strains | Fungal strains |
|                 |                 | SA   | EC   | CA   | CP   |
| V. guianensis   | EOVg            | >1000| >1000| >1000| 1.56 |
| V. cayennensis  | EOVc            | 25   | 50   | >1000| 50   |

*EOVg: Essential oil from Vismia guianensis; EOVc: Essential oil from V. cayennensis; SA: Staphylococcus aureus (ATCC 25923); EC: Escherichia coli (ATCC 25922); CA: Candida albicans (ATCC 10231); CP: Candida parapsilosis (ATCC 22019). Source: Authors.

OEVg inhibited fungal growth of C. parapsilosis strains at 1.56 µg/mL concentration. As stated by Holetz et al. (2002), samples with MIC values below 100 µg/mL are classified as very active, i.e., they strongly inhibit microbial growth. MIC values between 100 and 500 µg/mL denote satisfactory antimicrobial activity. Values between 500 and 1000 µg/mL indicate moderate activity whereas those above 1000 µg/mL imply poor inhibitory activity. In this study, OEVg exhibited strong antimicrobial activity against the strains under investigation.

Silvestre et al. (2012), investigating the essential oil extracted from V. guianensis fruit, reported antimicrobial activity against S. lentus (Kloos et al. 1976) Schleifer et al. 1983 (MIC equal to 78 µg/mL). Their research also identified β-caryophyllene (25.8%), α-copaene (13.1%), and δ-cadinene (11.6%) as the main components.

The main components of OEVg were (E)-caryophyllene sesquiterpenes (10.40%), α-copaene (29.45%), and (E)-nerolidol (24.06%). According to previous research, sesquiterpenes function as antimicrobial agents (CITÓ et al., 2003). The mechanism is still unclear, but it has been speculated that the lipophilic compounds found in this essential oil bind to and rupture the membrane of some microorganisms (COWAN, 1999). Reinsvold et al. (2011) showed that (E)-caryophyllene acts against microorganisms and, thus, can be used as antibiotic. This compound can also function biochemically as an anesthetic, anti-inflammatory, and spasmylytic drug. Other authors have reported antimicrobial activity of (E)-caryophyllene in kidney cell culture (RC-37), supporting its use as a prospective antimicrobial agent (Astani, 2009; Reichling & Schnitzler 2009).

Another study by Gelinski et al. (2007) indicates that (E)-nerolidol acts as a limited-spectrum antibiotic as it is not effective against some bacteria such as Salmonella Lignieres 1900 sp., E. coli, and Proteus Hauser 1885 sp. This finding
corroborates the results obtained in this study, i.e., the inactivity of OEVg against the bacteria *E. coli* and *S. aureus* as well as against the yeast *C. albicans*, probably due to this essential oil having this sesquiterpene as one of its main components.

The sesquiterpene α-copaene is one of the main components of copaiba (*Copaeifera L. sp.*). Studies have already shown that this oil can inhibit the growth of bacteria (Blose, 2003; Biavatti et al., 2006; Veiga & Pinto, 2002) and fungi (Craveiro et al., 1981; Wang, 2000; Souza et al., 2005). The microbial inhibitory capability of this oil is attributed to α-copaene.

OEVC exhibited strong inhibition against the bacteria *E. coli* and *S. aureus* at concentrations of 50 µg/mL and 25 µg/mL, respectively, and against the yeast *C. parapsilosis* at 50 µg/mL concentration. The main components identified in this essential oil are the oxygenated sesquiterpenes curzerene (25.29%) and germacrone (25.42%). Zhang and colleagues (2017) found curzerene to be one of the main components of the essential oil extracted from *Curcuma phaeocaulis* Valeton and attributed its observed antifungal activity (IC50, 153.33-580.09 µg/ml) and inhibition of bacterial growth (IC50, 485.00-778.33 µg/ml) to this sesquiterpene.

Another research conducted by Ogunwande et al. (2005) identified curzerene (19.7%) and germacrone (27.5%) as the main components of essential oils extracted from fruit and leaves of *Curcuma phaeocaulis*, respectively. The same study reports strong antibacterial activity for the essential oil extracted from *Eugenia uniflora* L. fruit and leaves against *S. aureus* and *Bacillus cereus*, respectively, with MIC equal to 39 µg/mL. In addition to treating cancer and hepatitis, other studies indicate that germacrone can be employed as an antimicrobial agent (Wang et al., 2000).

4. Conclusion

The findings of this study are relevant in that they show the antimicrobial potential of OEVg and EOVC against some microorganisms, namely, *C. parapsilosis* (EOVg and EOVC), *E. coli* (EOVC), and *S. aureus* (EOVC). This antimicrobial action can be explained by the presence, in large quantities, of some components in the plant species under investigation. The prevailing sesquiterpenes in EOVg — *E. coli* (E)-caryophyllene (10.40%), α-copaene (29.45%), and (E)-nerolidol (24.06%) — as well as the main components found in OEVc — *E. coli*, germacrone (25.42%) and curzerene (25.29%) — have demonstrated antimicrobial action. With the aim of addressing the paucity of information about essential oils from *V. guianensis* and *V. cayennensis*, this study has contributed their chemical profiles, indicated their antimicrobial potential, and, thus, provided a basis for future research with a view to isolating and characterizing the compounds responsible for the biological activities of OEVg and EOVC.

Acknowledgments

The authors would like to acknowledge Programa de Pós-Graduação em Ciência e Tecnologia para Recursos Amazônicos (PPGCTRA – UFAM), Fundação de Amparo à Pesquisa do Estado do Amazonas – FAPEAM (process nº. 001/2021 - Mulheres na Ciência). We are also grateful to the Herbarium from INPA and CESIT for the kindly support, Coordenação de Aperfeiçoamento de Pessoal de Nível Superior - Brasil (CAPES) and Conselho Nacional de Desenvolvimento Científico – CNPq.

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