Chemical Composition of Seasonal Essential Oils from *Psidium myrtoides* O. Berg Leaves with Antimicrobial, Antiprotozoal, Antioxidant and Anti-inflammatory Potential Activities

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Psidium species are traditionally used to treat bacterial agents and as an antiproliferative substance in human tumor cells. The current study investigated the seasonal essential oil composition of *Psidium myrtoides* O. Berg (Myrtaceae) leaves from the rainy and dry periods as well as some potential bioactivities. The essential oils were obtained by hydrodistillation using a Clevenger-type apparatus, with 2.0 L round-bottomed flask, with 100 g and 1.0 L of distilled water, at 97 °C, for 4 h. The obtained essential oils (EOs) were extracted with dichloromethane. The oil extraction procedure was performed in triplicate for each batch of leaves. The highest yield (1.4 ± 0.08%) was obtained in the rainy season oils. Twenty-six constituents were identified by gas-chromatography coupled to mass spectrometry (GC-MS) in the essential oils from the dry period and 22 constituents in the essential oils from the rainy season, all of them being hydrocarbons and oxygenated sesquiterpenes. The major compounds found in both periods, respectively, were β-caryophyllene (26.23%; 27.38%), α-humulene (17.40%; 16.18%), β-caryophyllene oxide (13.59%; 20.66%), humulene epoxide-II (6.09%; 10.21%), and α-copaene (8.92%; 7.11%). In general, no significant differences were observed in the chemical composition of essential oils, considering the seasonal effects and/or the status of the leaves. The antimicrobial and antiprotozoal activities were evaluated by the microdilution broth method and cytotoxic activity was carried out with Vero cells. The essential oils presented antibacterial activity against *Streptococcus sanguinis* (200 μg.mL⁻¹) and antifungal activity against *Candida glabrata* (46.87 μg.mL⁻¹) in Minimum Inhibitory Concentration (MIC) assays. The essential oil obtained from fresh leaves collected in the rainy season (FLRS) presented the highest Selectivity Index (0.23) against the studied promastigote forms. Therefore, the rainy volatile oils were more effective against promastigote forms of *Leishmania amazonensis* (31.8 ± 5.74 μg.mL⁻¹), while the dry volatile oils were more effective against *Trypanosoma cruzi* (54.0 ± 5.25 μg.mL⁻¹). Furthermore, this study investigated the antioxidant (ORAC method) capacity, using Trolox as the standard, and anti-inflammatory potential (*Drosophila melanogaster* of GMR-GAL4 and UAS-eiger lineage) of these oils. Thus, dry season contributed to a volatile oil (DLDS) with higher ORAC antioxidant capacity (2066.20 ± 168.20 μmol Trolox eq/g; p < 0.001) and showed no toxicity, as verified by the lineage survival test, being able to recover the eye area of the flies at 0.025-1.60 mg.mL⁻¹ (essential oils from the dry season) and at 0.2-1.6 mg.mL⁻¹ (essential oils from the rainy season) (p < 0.001). Therefore, the essential oils of *P. myrtoides* seems to be a potential therapeutical strategy against pathogens. In addition, our data suggest that the antioxidant and anti-inflammatory activities of the studied essential oils may be, at least in part, due to donation of hydrogen atoms and suppression of the Eiger pro-inflammatory cytokine, respectively.

**Keywords:** *Psidium myrtoides*; essential oil; terpenes; bioactivities assays

1. Introduction

The Myrtaceae family is generally considered as an important edible fruit family, which comprises around 121 genera and 3800-5800 species of woody shrubs or trees, mostly evergreen and distributed mainly in the tropics and subtropics.¹ *Psidium myrtoides* O. Berg (Myrtaceae) is popularly known as “araçá-roxo” in the Cerrado biome, Brazil, having the appearance of trees of 6 m to 12 m in length and, sometimes reaching 20 m. The bark of the trunk is exfoliating with papyraceous blades, and it presents opposite leaves with a size between 2.5 cm and 7.5 cm. Its flower bearing bud opens with 4 flowers. The fruits, when ripe, present an...
atropurpureum color and are edible. The pharmacological properties of araçá species are attributed to their fruits and leaves, such as antioxidant, antiprotozoal, antibacterial, and antiproliferative effect in human tumor cells. However, its antioxidant and anti-inflammatory potential have not been explored yet.

The essential oils extracted from leaves of the Psidium genus are rich in terpene compounds. The β-caryophyllene is the most common (36.7%) sesquiterpene, especially in the P. guajava and P. parvifolium species. Other terpene compounds usually present are α-pinene (12.9%), 1,8-cineole (12.3%), (β)-caryophyllene oxide (12.0%), limonene (6.4%), α-humulene (4.4%), α-terpineol (3.8%), β-cadinene (3.7%), and β-selinene (3.5%). The rate of sesquiterpenes in the genus Psidium is approximately 32%, and most of these chemical compounds are obtained from the leaves (79%). These compounds have well-described antioxidant and anti-inflammatory properties and may provide the Psidium myrtoides O. Berg the ability to scavange the reactive oxygen species and to modulate the inflammation processes.

In this scenario, the transgenic lineage of Drosophila melanogaster, the GMR-GAL4>UAS-eiger lineage, has been standardized as an animal model for evaluating the anti-inflammatory potential of different new molecules and natural products. The flies of this strain have an anti-inflammatory potential using the GMR-GAL4>UAS-eiger lineage. Therefore, this study aimed to elucidate the seasonal chemical composition of the essential oils obtained from the leaves of Psidium myrtoides, comparing them according to the period of collection (dry and rainy seasons), and to evaluate their in vitro antimicrobial, antiprotozoal, antioxidant, and anti-inflammatory potential using the D. melanogaster of the GMR-GAL4>UAS-eiger lineage.

2. Materials and Methods

2.1. Plant materials

The undamaged leaves of Psidium myrtoides were randomly collected from the top of single adult tree in the local area of Uberlândia (18°56’45.3" S and 48°16’08.4" W), Minas Gerais, Brazil. The collection was carried out in two different periods, in September 2017 (11:00 a.m.; dry season; without rainfall) and February 2018 (11:00 a.m., rainy season; average of 125 mm rainfall). There was no separation between young and old leaves. The dry leaves were dried at room temperature for 72 h.

The authorization for collecting the plant was issued by the CNPq/SISGEN under the code A1247B0, and a voucher specimen is registered in the herbarium Uberlândense with the collection code HUFU65831.

2.2. Essential oil extraction

The extraction of the volatile oil using fresh or dried leaves was done by hydrodistillation using a Clever-type apparatus with 2.0 L round-bottomed flask, with 100 g of crushed samples and 1.0 L of distilled water, at 97 °C, for 4 h. The oil extraction procedure was performed in triplicate for each batch of leaves. The obtained essential oils (EOs) were extracted with dichloromethane; the organic phase was separated and dried with anhydrous sodium sulphate; filtered and kept in a closed vial under refrigeration (-10 °C) for further analysis. The percentage yield was calculated based on the dry mass of the initial sample, with this procedure being performed in triplicate.

2.3. Analysis and identification of the constituents

The oils were analyzed, and their constituents were identified by gas-chromatography coupled to mass spectrometry (Shimadzu, model GC17A/QP5010), equipped with a SP-5 capillary column 30 m × 0.25 mm × 0.25 μm film thickness). The GC-MS chromatograms for essential oils from the dry and rainy periods are shown in Figures 1S and 2S (Supplementary Material). The carrier gas was helium at a flow rate of 1 mL.min⁻¹, detector and injector temperatures were 220 °C and 246 °C respectively, the injection volume was 1 μL, and the split ratio was 1:20. The oven temperature was programmed from 60 °C to 246 °C, at 3 °C min⁻¹ and the injected volume was 1 μL of 5 μg.mL⁻¹ solution in dichloromethane.

The electron impact energy was set at 70 eV and fragments from 40 m/z to 650 m/z were collected. The identification of the essential oil components was carried out by matching the spectra obtained with those stored in software libraries (Wiley7; Wiley229; Nist08; Nist08s; Nist27; Shim2205) and by comparing the calculated arithmetic index (AI) with the arithmetic index reported in the literature. Only peaks higher than with a match factors above 90% were considered. Authentic standards were used when necessary. Arithmetic indices were calculated using equation $AI (X) = 100 \frac{PzC}{C - C_0} + 100\left(\frac{t(X) - t(Pz)}{t(Pz+1) - t(Pz)}\right)$, based on the retention times (R) of linear alkane standards (C₂-C₇0) which, by definition, have an AI equal to 100 times the number of carbon atoms. X = compound at time t; PzC = number of carbon atoms of the alkane Pz, which runs just before X; Pz+1 = alkane running after X. Quantification was obtained after normalization of the peak areas in the total ion chromatogram (TIC). Results represent average values of three replicates for each essential oil analysis.
Only concentrations over 1.0% (TIC-area) were considered. The identification was based on a comparison of the mass spectra obtained and those of the virtual libraries, and the AI obtained was compared with AIs of the NIST Standard Reference Data.12

2.4. Anticandidal assay

The microorganisms tested were Candida albicans (C. albicans) (ATCC 28366), Candida tropicalis (C. tropicalis) (ATCC 13803), and Candida glabrata (C. glabrata) (ATCC 15126) from the American Type Culture Collection (ATCC, Rockville MD®, USA). The assays were performed via the broth microdilution method using the standards recommended by the Clinical and Laboratory Standards Institute.13

The stock solutions were prepared by dissolving the essential oils in dimethyl sulfoxide (DMSO) (Sigma-Aldrich, Co) in a concentration of 192,000 μg.mL⁻¹. Dilutions were made using the standard RPMI 1640 medium, at 12,000 μg.mL⁻¹. The inoculum was prepared using a spectrophotometric method (at 530 nm wavelength) and compared with a 0.5 McFarland scale to obtain 6.0 × 10⁶ CFU mL⁻¹. Then, the dilutions recommended by the CLSI were made with RPMI until the inoculums reached 1.2 × 10³ CFU/mL.¹³

The minimum inhibitory concentration (MIC) was determined using the broth microplates, where dilutions with final concentrations were made in the range from 1.46 to 3,000 μg.mL⁻¹. Amphotericin B was used as positive control, being diluted in broth at concentrations between 0.031 and 16.0 μg.mL⁻¹. The negative control (DMSO) was dissolved in dimethyl sulfoxide (DMSO, Sigma–Aldrich, St. Louis, USA) at 8000 μg.mL⁻¹, followed by a dilution in tryptic soy broth (TSB; Merck, Darmstadt, Germany) for aerobic microorganisms; and in Schaedler broth (BBL, Microbiology Systems, Cockeysville, MD) supplemented with hemin (5.0 μg.mL⁻¹) and vitamin K₁ (10.0 μg.mL⁻¹), for anaerobic microorganisms. The concentrations tested ranged from 400 μg.mL⁻¹ to 25 μg.mL⁻¹. The final DMSO content was 4% (v/v), and this solution was used as a negative control. The inoculum was adjusted for each organism to yield a cell concentration of 5 × 10⁷ colony forming units (CFU) per mL. The microplates with the anaerobic microorganisms were incubated aerobically at 37 °C, for 24 h. The aerobic microorganisms were incubated for 48-72 h in an anaerobic chamber (Don Whitley Scientific Bradford, UK), at 5 °C. After that, resazurin (Acros Organics NV, Geel, Belgium) in aqueous solution (0.01% w v⁻¹) was added to the microplates to indicate the microorganism viability. Chlorhexidine dihydrochloride was used as a positive control. The negative control (DMSO) was tested with concentrations ranging from 1% to 10% (v v⁻¹) which did not affect the bacterial growth. Sterility tests were performed for the TSB and Schaedler broths, control culture (inoculum), positive control, oils, and DMSO.

2.6. Antiprotozoal activity

The essential oils from P. myrtoides were dissolved in methanol and diluted with Dulbecco’s Modified Eagle Medium (DMEM) to test their efficacy against Trypanosoma cruzi (T. cruzi) and diluted with BHI broth to evaluate their properties against Leishmania amazonensis (L. amazonensis) to a final stock solution of 1280 μg.mL⁻¹ was obtained.¹³ The cell viability test was carried out, using tissue-culture tryptomastigote (G strain) and a L. amazonensis promastigote stage (PH8 strain). The analysis was performed using a microdilution plate of 96 wells, obtained from the stock solution at concentrations of 1024, 512, 256, 128, 64, 32, 16, and 8 μg.mL⁻¹, not exceeding a value of 3% for methanol concentration. The final volume in each well was 100 μL: 20 μL of inoculum (1 × 10⁷ parasites in 2 mL) and 80 μL of sample. Positive, negative, and control samples were prepared on each plate. The T. cruzi plate was incubated, for 48 h, at 37 °C, in a humidified atmosphere with 5% CO₂. Then, 4 μL of 3 mM resazurin in phosphate buffer solution (PBS) were added to each well as a developing solution, and incubated for 24 h under the same conditions, as outlined above. The L. amazonensis plate was incubated for 48 h, at 25 °C, and then 2 μL of developing solution of 3 mM resazurin in PBS was added to each well and incubated again for 24 h, at 25 °C. At the
end of this period, the absorbance (594 nm) was read using a microplate spectrophotometer. The half-maximal inhibitory concentration (IC$_{50}$) was calculated in triplicate using a dose-response graph with non-linear regression for each test.

2.7. Cytotoxicity

This test followed a procedure adapted from Martins et al.,$^{14}$ with the essential oil samples of $P$. myrtoides being dissolved in methanol, diluted in DMEM (Sigma Aldrich, USA), and supplemented until a solution of 1280 μg.mL$^{-1}$ was achieved. The cell viability test was performed with ATCC Vero cells (CCL 81; kidney fibroblasts from the African green monkey). The microplate dilution method was used for cytotoxicity evaluation. In each test, a solution was prepared containing $1 \times 10^6$ cells in supplemented DMEM. Then, 100 μL of this solution was added into each well, and the plate was incubated for 6 h, at 37 °C, in a humidified atmosphere with 5% CO$_2$, causing adhesion to the well. Once cells were attached, the culture medium was removed and solutions of the samples were added, reaching the concentrations of 1,024, 512, 256, 128, 64, 32, 16, and 8 μg.mL$^{-1}$ by serial dilutions from the stock solution. The final concentration of methanol in each well did not exceed 3%. Control negative (100% lysed cells), solvent (methanol), and samples were also prepared. The microplates were incubated for 48 h, at 37 °C, in a humidified atmosphere with 5% CO$_2$. Afterwards, a revealing solution of resazurin 3 mM in PBS (Sigma Aldrich, St. Louis, Missouri, USA) was added to each well, and the plate was incubated again, for 24 h, under the same conditions. Absorbance was read at 594 nm using a microplate spectrophotometer (Spectra Max190, Molecular Devices). The assays were performed in five replicates and the results of absorbance for each concentration tested were calculated according to the growth in the control. The cytotoxic concentration (CC$_{50}$) at which 50% of the cells are viable was calculated using a dose-response graph with non-linear regression for each test. The relationship between CC$_{50}$ from the Vero cell line and IC$_{50}$ from $L$. amazonensis was established by the Selectivity Index (SI) and calculated according to the equation proposed by Case et al.$^{15}$

2.8. Oxygen radical absorbance capacity (ORAC) antioxidant activity test

The ORAC method was done according to Justino et al. (2018).$^{16}$ The essential oils of $P$. myrtoides diluted in methanol (1 μg.mL$^{-1}$) were incubated with 0.085 nM fluorescein (diluted in 75 mM sodium phosphate buffer, pH 7.4), at room temperature, for 15 min. The mixture received 2,2’-azobis (2-amidino-propane) dihydrochloride (153 mM, diluted in 75 mM sodium phosphate buffer, pH 7.4) and the loss of fluorescence intensity (485 nmex/528 nmex) (Perkin-Elmer LS 55, Waltham, MA, USA) was measured every 2 min, for 90 min, at 37 °C, by calculating the area under the curve. The antioxidant capacity was determined using an analytical curve, constructed using Trolox as the standard (μmol trolox equivalents/g). Ascorbic acid was used as positive control and all analyses were done in triplicate.

2.9. Anti-inflammatory potential assay

2.9.1. Acquisition of flies

The transgenic strain of $Drosophila$ melanogaster GMR-GAL4 (driver) > UAS-eiger (reporter) was obtained from Bloomington (Drosophila Stock Center – Indiana University), kept in the Genetics Laboratory of the Federal University of Uberlândia, kindly provided by professor Dr. Carlos Ueira. The drosophila host were kept in glass flasks containing a culture medium prepared with water (150 mL), yeast (2.7 g), soy flour (1.5 g), cornmeal (10.95 g), agar (0.9 g), glucose syrup (11.55 g), acid solution (1.25 mL), and Nipagin solution (1.5 mL), remaining in a BOD oven (SOLAB, Piracicaba, SP, Brazil), in a 12:12-h light/dark (12L:12D) photoperiods, at 25 °C.

2.9.2. Survival and inflammatory assays

The survival and inflammatory assays with GMR-GAL4>UAS-eiger were based on the model proposed by Igaki et al. (2002),$^9$ with modifications, performed on larvae in the L3 stage of development. Briefly, adult flies were transferred to a laying medium composed of fresh biological yeast supplemented with sucrose, for 8 h, to perform oviposition. After 72 h, the new individuals in the L3 stage were collected and transferred to a culture medium containing 2 g of mashed potato (Yoki, Brazil), and 5 mL of the essential oils of $P$. myrtoides leaves collected in the dry season or in the rainy season (0.025 - 1.60 mg.mL$^{-1}$, diluted in 0.025% Tween 80). The treatments with water (negative control) or vehicle (0.025% Tween 80) were also tested. Each treatment used 40 larvae in the L3 stage. The development of the flies was monitored, and the number of individuals that reached the pupal and adult phases was recorded. The adult flies that hatched in the groups treated with essential oils and, in the control groups, were anesthetized with ether and fixed in 70% ethanol to preserve the morphology of the organisms, which were subsequently photographed with a stereoscopic microscope (Nikon SMZ 800), with 3X zoom, coupled to a digital camera and using the IC Capture software. The eyes of each fly were photographed in right and left views, and the eye area, defined as the red area in the ocular region of the individual was selected and measured using the ImageJ software.

2.10. Statistical analysis

The statistical analyses and graphics were done using GraphPad Prism 6.0 software (GraphPad Software Inc., San Diego, CA, USA). The data were expressed as mean ± standard error and the statistical significance of the difference was calculated using the unpaired t-test and one-
The concentrations of β-caryophyllene and β-caryophyllene oxide are quite similar in the essential oil from fresh leaves collected in the rainy season (18.02% and 20.66%, respectively), but when the leaves were dried for 72 h the amount of β-caryophyllene oxide dropped dramatically (from 20.66% to 11.41%). The major compound in all essential oils was β-caryophyllene, probably because of the direct sunlight exposure of the plant in a tropical region, and due to the greater humidity in the fresh leaves (50% versus 16% in the dried leaves). This result can also be related to the elimination of substances when the leaves are dried, making some compounds more evident than the others, to the detriment of their availability on the surface or inside the leaf along with the higher solar exposure of leaves.

3.2. Antifungal and antibacterial activities

The Candida species evaluated in this work are often involved with opportunistic infections, with 95% of the bloodstream infections being caused by Candida species such as C. albicans, C. tropicalis, C. glabrata, and C. parapsilosis. These yeasts were resistant to at least one of the following drugs amphotericin B, fluconazole, and itraconazole. The MFC found for C. glabrata in the present study is significant since this yeast from the hospital environment presented one of the greatest resistances against fluconazole (64.7%), according to Furlaneto et al. Others studies have shown that about 10% of C. glabrata present resistance to fluconazole, and about 52% are resistant to itraconazole. The EO of Blepharocalyx salicifolius, as example, with high sesquiterpene hydrocarbon content, was tested against different species of Candida and did not show activity in concentrations up to 5,000 μg.mL-1. β-caryophyllene oxide and β-caryophyllene tested alone showed weak activity for C. glabrata and C. tropicalis and were inactive for C. albicans (> 3,000 μg.mL-1).

It is well known that essential oils rich in oxygenated terpenes can inhibit the development of Candida biofilms. Among the compounds found in the essential oil from P. myrtoides, β-caryophyllene oxide (11.41%-13.59%), β-bisabolol (0.89%-2.65%), and α-copaene (4.86%-8.92%) presented some antifungal action in synergism with other oxygenated terpenes.

According to the literature, the antifungal and antibacterial activities of essential oils is classified as strong (up to 500 μg.mL-1), moderate (between 500 and 1500 μg.mL-1), and weak (higher than 1500 μg.mL-1). The essential oil from dried leaves collected in the dry season showed strong activity for C. glabrata, with a MIC of 46.9 μg.mL-1, while the essential oils from fresh leaves sampled in the rainy and dry seasons, and dried leaves collected in the rainy season presented a strong effect against this microorganism, with MICs ranging from 187.5 μg mL-1 to 375.0 μg.mL-1 (Table 2). The antifungal activities for C. tropicalis were moderate to weak with MICs ranging
Table 1. Chemical composition of the essential oils from fresh and dried leaves of *Psidium myrtoides* O. Berg from the dry and rainy seasons

| Peak No | Compounds dry season | Peak No | Compounds rainy season | AI literature (on OV-5) | FLDS | DLDS | FLRS | DLRS | Composition (% TIC) |
|---------|----------------------|---------|------------------------|-------------------------|------|------|------|------|---------------------|
| 1       | α-Copaene            | 1       | α-Copaene              | 1377                    | 1383 | 1379 | 1334 | 1382 | 8.92 (6.97)         |
| 2       | β-Caryophyllene      | 2       | β-Caryophyllene        | 1417                    | 1430 | 1423 | 1375 | 1427 | 26.23 (21.62)       |
| 3       | Aromadendrene        | 3       | trans-α-Bergamoptene   | 1392                    | -    | -    | 1443 | 1427 | - (1.31)            |
| 4       | α-Humulene           | 4       | Aromadendrene          | 1439                    | 1443 | 1442 | 1408 | 1446 | 1.40 (1.34)         |
| 5       | Ishwarane            | 5       | α-Humulene             | 1452                    | 1446 | 1458 | 1414 | 1462 | 15.56 (17.40)       |
| 6       | δ-Selinene           | 6       | Ishwarane              | 1465                    | 1464 | 1464 | 1431 | 1465 | 0.78 (0.93)         |
| 7       | trans-α-Bergamoptene | 7       | δ-Selinene             | 1492                    | 1495 | 1499 | 1440 | 1495 | 1.04 (1.79)         |
| 8       | trans-α-Cadinene     | 8       | δ-Selinene             | 1498                    | 1495 | 1497 | 1449 | 1500 | 1.06 (2.70)         |
| 9       | trans-Cadin-1,4-diene| 9       | δ-Cadinene             | 1513                    | 1499 | 1519 | 1464 | 1515 | 2.82 (0.76)         |
| 10      | α-Cadinene           | 10      | trans-Cadin-1,4-diene  | 1533                    | 1519 | 1528 | 1465 | 1529 | 0.92 (2.14)         |
| 11      | Elemol               | 11      | α-Cadinene             | 1537                    | 1534 | 1539 | 1477 | 1540 | 4.76 (1.66)         |
| 12      | (E)-Nerolidol        | 12      | Elemol                 | 1548                    | 1544 | 1545 | 1487 | 1547 | 1.49 (1.75)         |
| 13      | Spathulenol          | 13      | (E)-Nerolidol          | 1561                    | 1551 | 1561 | 1563 | 1561 | 1.99 (2.28)         |
| 14      | β-Caryophyllene oxide| 14      | Spathulenol            | 1577                    | 1566 | 1569 | 1508 | 1570 | 2.51 (1.20)         |
| 15      | Globulol             | 15      | β-Caryophyllene oxide  | 1582                    | 1574 | 1587 | 1518 | 1590 | 7.69 (13.59)        |
| 16      | Humulene epoxide II  | 16      | Globulol               | 1608                    | 1593 | 1607 | -    | -    | 0.65 (0.35)         |
| 17      | Cubenol              | 17      | Humulene epoxide II    | 1608                    | 1620 | 1613 | 1533 | 1617 | 2.61 (6.09)         |
| 18      | Muurola-4,10(14)-dien-1β-ol | 18   | Cubenol               | 1645                    | 1627 | 1634 | -    | -    | 3.62 (3.79)         |
| 19      | Caryophylla 4(12), 8(13)-dien-5β-ol | 19 | Muurola-4,10(14)-dien-1β-ol | 1630                    | 1640 | 1639 | 1557 | 1636 | 0.33 (0.22)         |
| 20      | Selina-3,11-dien-6α-ol| 20      | Caryophylla 4(12), 8(13)-dien-5β-ol | 1639                    | 1646 | 1643 | -    | -    | 0.12 (0.24)         |
| 21      | α-Muurolol           | 21      | Selina-3,11-dien-6α-ol | 1642                    | 1658 | 1651 | -    | -    | 0.03 (0.29)         |
| 22      | Neo-Intermedeol      | 22      | α-Muurolol            | 1644                    | 1669 | 1653 | 1577 | 1655 | 1.79 (1.50)         |
| 23      | N.I.                 | 23      | Neo-Intermedeol       | 1658                    | 1672 | 1662 | 1590 | 1663 | 1.33 (1.17)         |
| 24      | β-Bisabolol          | 24      | N.I.                 | 1683                    | 1683 | 1651 | -    | -    | 0.51 (1.31)         |
| 25      | α-Bisabolol          | 25      | β-Bisabolol          | 1674                    | 1687 | 1677 | 1601 | 1679 | 0.89 (2.23)         |
| 26      | Eudesm-7(11)-en-4α-ol| 26      | α-Bisabolol          | 1685                    | 1701 | 1693 | 1618 | 1694 | 10.16 (5.89)        |

Functional group

| Functional group | % TIC (n. of compounds) |
|------------------|-------------------------|
| Sesquiterpenes hydrocarbons | 63.50 (10) |
|                   | 57.00 (10)  |
|                   | 47.07 (11)  |
|                   | 62.79 (11)  |
| Sesquiterpenes Oxygenated | 35.99 (15) |
|                   | 41.62 (15)  |
|                   | 52.88 (11)  |
|                   | 37.21 (11)  |
| N.I.              | 0.51 (1)    |
|                   | 1.01 (1)    |
| TOTAL (% TIC: n. compounds) | 100.00 (26) |
|                   | 99.93 (26)  |
|                   | 99.95 (22)  |
|                   | 100.00 (22) |

FLDS: fresh leaves essential oil from the dry season; DLDS: dried leaves essential oil from the dry season; \(^1\) Arithmetic indices from the literature (ref.5); \(^2\) Arithmetic indices determined in relation to the standard \(n\)-alkanes (C\(_8\) - C\(_{30}\)) in column OV-5 (30 m X 0.25; 025 μm); TIC = total ions chromatogram, by three replicates. N.I. = Unidentified. FLRS: fresh leaves essential oil from the rainy season; DLRS: dried leaves essential oil from the rainy season.
from 375 µg.mL⁻¹ to 3,000 µg.mL⁻¹, while for *C. albicans* the leaves had a weak effect on fungus, with MICs ranging from 3,000 µg.mL⁻¹ to 1,500 µg.mL⁻¹.

Regarding antibacterial activity the essential oils from fresh and dried leaves sampled in the dry season (FLDS and DLDS) showed moderate activity against *S. mutans* (400 µg.mL⁻¹) and *S. sanguinis* (200 µg.mL⁻¹). The essential oils from fresh and dried leaves collected in the rainy season (FLRS and DLRS) were active only against *S. sanguinis* (200 µg.mL⁻¹) and none of the oils had any effect on *S. mitis* (Table 2).

### 3.3. Antiprotozoal and cytotoxicity activities of the essential oils

Antiprotozoal activity (IC₅₀) and cytotoxicity (CC₅₀) data were obtained from the cell viability tests. According to the results of Table 3, essential oils from fresh and dried leaves collected in the dry season presented the lowest IC₅₀ values of 796.30 ± 8.93 µg.mL⁻¹ and 54.05 ± 5.25 µg.mL⁻¹, respectively, for *Trypanosoma cruzi*. This indicates that for the DLDS a higher concentration is required to inhibit 50% of parasites.

Based on the in vitro assay results, it was possible to calculate the concentration that determined 50% of cell viability (CC₅₀) and relate it to the IC₅₀ values for the calculation of the Selectivity Index (SI). A positive value represents higher selectivity against microorganisms than cellular toxicity, and a negative value, lower selectivity for microorganisms and higher cytotoxicity. Thus, FLDS is more toxic (IC₅₀ 796.3 ± 8.93) than selective for the parasite with negative selectivity index (SI) values.

Regarding the essential oils from fresh and dried leaves sampled in the rainy season, there was no inhibition against *T. cruzi* at the concentration tested. In the *L. amazonensis* assay, essential oils from fresh and dried leaves collected in the dry season presented no inhibitory activity against the

### Table 2. Inhibitory effect on the growth of yeasts, aerobic, and anaerobic oral bacteria (MIC values, µg.mL⁻¹) of the essential oils from *P. myrtoides* leaves collected in the dry and rainy seasons

| Microorganism          | Samples | MIC (µg.mL⁻¹) | FLDS | DLDS | FLRS | DLRS | OC | TC | Positive control |
|------------------------|---------|---------------|------|------|------|------|----|----|------------------|
| **Gram-positive**       |         |               |      |      |      |      |    |    |                  |
| *Streptococcus mutans* |         |               | 400  | 400  | > 400| > 400| > 400| > 400| 0.92a            |
| *Streptococcus mitis*  |         |               | > 400| > 400| > 400| > 400| > 400| > 400| 3.68a            |
| *Streptococcus sanguinis* |       |               | 200  | 200  | 200  | > 400| > 400| > 400| 0.92a            |
| *A. actinomycetencomitans* |     |               | > 400| > 400| > 400| > 400| > 400| > 400| 0.46a            |
| *Actinomyces naeslundii*|         |               | > 400| > 400| > 400| > 400| > 400| > 400| 1.84a            |
| *Staphylococcus aureus* |         |               | > 400| > 400| -    | -    | > 400| > 400| 6.75a            |
| **Gram-negative**       |         |               |      |      |      |      |    |    |                  |
| *Escherichia coli*      |         |               | > 400| > 400| -    | -    | > 400| > 400| 3.12a            |
| *Pseudomonas aeruginosa*|        |               | > 400| > 400| -    | -    | > 400| > 400| 12.5a            |
| *Porphyromonas gingivalis* |       |               | > 400| > 400| > 400| > 400| > 400| > 400| 3.68a            |
| *Fusobacterium nucleatium* |     |               | > 400| > 400| > 400| > 400| > 400| > 400| 1.84a            |
| **Yeast**               |         |               |      |      |      |      |    |    |                  |
| *Candida albicans*      |         |               | 3000 | 3000 | 1500 | 1500 | > 3000| > 3000| 0.25c            |
| *Candida glabrata*      |         |               | 187.5| 46.87| 187.5| 375  | 750  | > 3000| 0.25c            |
| *Candida tropicalis*    |         |               | 1500 | 1500 | 375  | 750  | > 3000| 750   | 0.12c            |

FLDS: fresh leaves essential oil from the dry season; DLDS: dried leaves essential oil from the dry season; FLRS: fresh leaves essential oil from the rainy season; DLRS: dried leaves essential oil from the rainy season. Positive control: ¹ Chlorhexidine dihydrochloride; ² Chloramphenicol; ³ Anfhotericin B. CO: β-Caryophyllene oxide. TC: β-caryophyllene. Bacteria control by protocol M11-A8 CLSI (2012): *B. fragilis* (MIC 0.73 µg mL⁻¹); *B. theitaiotaomicron* (MIC 1.47 µg mL⁻¹); Yeast control by protocol M27-A3 CLSI (2008): *C. krusei* (MIC 1.0 µg mL⁻¹); *C. parapsilosis* (MIC 0.5 µg mL⁻¹)
parasite at the concentration tested. Essential oils from fresh and dried leaves sampled in the rainy season presented low IC₅₀ values. The results obtained for all volatile oils tested were correlated by calculating the SI (Table 3). Based on the in vitro assay results, it was possible to calculate the concentration that determined 50% of cell viability (CC₅₀), and to relate it with IC₅₀ values for the calculation of the selectivity index (SI). A positive value represents higher selectivity against microorganisms than cellular toxicity, and a negative value, lower selectivity for microorganisms and higher cytotoxicity.¹⁵

Regarding antiprotozoal activity (IC₅₀), the essential oils from fresh and dried leaves from the dry season presented inhibitory activities against trypomastigote forms of *T. cruzi* but not to the promastigote forms of *L. amazonensis*. The opposite occurred with the essential oils from rainy season leaves (Table 3). According to Chibale et al.,²⁸ the lower the IC₅₀ value (inhibitory concentration of 50% of the protozoa) the higher was the antiprotozoal activity. The results in this study showed that the volatile oils from fresh leaves of *P. myrtoides* O. Berg collected in the rainy season were the most effective against the promastigote forms of *L. amazonensis* (31.8 ± 5.74 µg.mL⁻¹), followed by dried leaves from the same season (55.7 ± 7.86 µg.mL⁻¹). The antiprotozoal activity of the essential oils depended on the type of terpenes, since they determine the interaction between the volatile oils with the cell hydrophobic membrane of the protozoa.²⁹ However, some controversies persist regarding the efficacy of terpenes on the antiprotozoal activity. While monoterpenes are more effective than sesquiterpenes against the forms of *L. amazonensis* ²⁹,³⁰ the inhibition of protozoa results from the action of sesquiterpenes (i.e. α-bisabolol). Other sesquiterpene compounds exert trypanosomicidal activity in vivo and in vitro, like (E)-nerolidol ¹³ and β-caryophyllene ¹³,³⁰,³² compounds also found in this study.

The cytotoxicity test, using Vero cells, revealed that all oils were toxic to the cells, which means that they contained few promising compounds for isolation and to be used against *T. cruzi* and *L. amazonensis* (Table 3). Moreover, positive SI values were found in the oils obtained from dried leaves sampled in the dry season as well as from dried and fresh leaves collected in the rainy season, representing a higher selectivity against microorganisms and low toxicity to Vero cells lineage. Negative SI values found in the fresh leaves collected in the dry season presented a higher toxicity to Vero cells and lower selectivity to the trypomastigote forms.¹⁴ Despite the low SI values, the oils were selective to inhibit the parasite. Within this context, the oils from dried leaves sampled in the dry season as well as from dried and fresh leaves from the rainy season could be a promising agent for the treatment of leishmaniasis and Chagas disease.

Although these studies have been done with the same species, this activity difference may be explained by qualitative and quantitative variations in the chemical composition of the oils, as well as the consequence of seasonality and different collection regions.³³,³⁴

### 3.4. In vitro total antioxidant capacity of the essential oils from the leaves of *P. myrtoides*

The essential oils from leaves collected both in the dry and rainy seasons presented antioxidant properties according to the ORAC method (Table 4). The dry season contributed to a higher antioxidant capacity of the essential oils when compared with the rainy season (*p* < 0.001).

The antiradical activity of essential oils is mainly based on the hydrogen-donating properties of terpenes, corroborating the ORAC results for the essential oils of *P. myrtoides*. Furthermore, as expected, the essential oils extracted from leaves collected in the dry season showed a

| Sample       | Cytotoxicity (Vero cells) CC₅₀ µg.mL⁻¹ | Trypanosoma cruzi IC₅₀ µg.mL⁻¹ | Leishmania amazonensis IC₅₀ µg.mL⁻¹ | SI       |
|--------------|--------------------------------------|-------------------------------|-----------------------------------|----------|
| FLDS-EO      | 58.99 ± 10.41                        | 796.30 ± 8.93                 | No toxic                          | -1.13    |
| DLDS-EO      | 71.77 ± 6.41                         | 54.05 ± 5.25                  | No toxic                          | 0.12     |
| FLRS-EO      | 53.79 ± 6.55                         | No toxic                      | 31.80 ± 5.74                      | 0.23     |
| DLRS-EO      | 66.57 ± 1.69                         | No toxic                      | 55.70 ± 7.86                      | 0.08     |

FLDS-EO: fresh leaves essential oil from the dry season; DLDS-EO: dried leaves essential oil from dry season; FLRS-EO: fresh leaves essential oil from the rainy season; DLRS-EO: dried leaves essential oil from rainy season

| Antioxidant capacity | Essential oils |
|----------------------|---------------|
| ORAC (µmol Trolox eq/g) | DLDS | DLRS |
| 2066.20 ± 168.20 | 1144.90 ± 38.30 |

*Values (mean ± standard error) are expressed as µmol Trolox equivalents/g for ORAC assay. Unpaired t-test, *p* < 0.001. DLDS = dried leaves from dried season; DLRS = dried leaves from rainy season."
higher antioxidant ability, which may be explained by the higher concentration of terpenes when compared with the essential oils from rainy season leaves.5,6,7

3.5. Anti-inflammatory potential of the essential oils from the leaves of *P. myrtoides*

The use of plants with medicinal properties as antioxidant and anti-inflammatory agents has gained prominence in the control and treatment of acute and chronic inflammatory conditions. Plant-derived terpenes, specifically sesquiterpenes, have anti-inflammatory effects, with suggested modulation of the TNF-α and nuclear transcription factor-kappa B (NF-κB) signaling pathway.8 Our results demonstrated that the cell death activity induced by the overexpression of the eiger gene (TNF-α) in the eyes of the flies was reduced through the chronic exposure of L3 larvae to the essential oils, showing that the treatment has the ability to inhibit the activation of the induced c-Jun N-terminal kinase pathway by the pro-inflammatory cytokine.9,10 The treatments with the essential oils from *P. myrtoides* leaves sampled in the dry and rainy seasons, as well as with water and vehicle, were carried out with 40 L3 larvae, with the survival rate of the flies shown in the Figure 2.

Although the essential oils from leaves collected in the dry season showed low toxicity for the pupae and adult flies at 1.6 mg.mL⁻¹, there was no change in the survival pattern of the pupae and adult flies treated with the essential oils, only with water, or with the vehicle (Figures 2a and 2b).

This study also revealed the *in vitro* antioxidant properties of the *P. myrtoides* dried leaves as well as the ability to recover the eye area of a transgenic strain of Drosophila melanogaster.16,15-38

Studies on medicinal plants have shown a correlation between the antioxidant and anti-inflammatory properties.39,40 Therefore, this study investigated the anti-inflammatory potential of the essential oils in a transgenic strain of *D. melanogaster* GMR-GAL4> UAS-Eiger. This fly presents a cytokine that is orthologous to the pro-inflammatory cytokine TNF-α (Eiger in *D. melanogaster*) and its respective receptor.41

TNF-α is a key cytokine in different inflammatory diseases and there are clinical therapies focused on its

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**Figure 2.** Survival of the transgenic strain of *Drosophila melanogaster* GMR-GAL4> UAS-eiger submitted to a treatment with essential oils of *Psidium myrtoides* extracted from dried leaves collected in the dry (a) and rainy (b) seasons (0.025 – 1.6 mg mL⁻¹) and quantification of the eye area of the flies treated with essential oils of *P. myrtoides* extracted from dried leaves from the dry (c) and rainy (d) seasons (0.025 – 1.6 mg mL⁻¹). Columns represent the mean ± standard error. Note: -: control treatment with water; V: treatment with vehicle (0.025% Tween 80). One-way ANOVA followed by Dunnett as post-test, ***p < 0.001 compared with the control group (-). Note: -: control treatment with water; V: treatment with vehicle (0.025% Tween 80)
control. Additionally, the use of fruit flies as Drosophila for research is in accordance with the 3R’s rule of reduction, replacement, and refinement, making this an excellent biological model for inflammation analysis. In accordance with our hypothesis, a partial recovery of the eye area was observed in adult flies treated with the P. myrtoides essential oils compared to untreated flies, both for oil extracted during the dry and rainy periods, indicating the inhibition of the eiger gene’s action (Figure 3S, Supplementary Material).

The flies treated only with water or vehicle (Figures 3S-a and 3S-b, respectively) did not present ocular recovery. However, the treatment with all concentrations of the essential oils from leaves sampled in the dry season (Figure 3S-c) presented an increase in the eye area (p < 0.001) when compared with the treatment with the control (water) or vehicle. Regarding the essential oils collected in the rainy season, concentrations above 0.2 mg.mL\(^{-1}\) induced an increase in the eye area of the flies (Figure 3S-d) (p < 0.001).

As expected, and in accordance with the antioxidant test results, the eye recovery effect was observed for all concentrations tested of the essential oils from leaves collected in the dry season (0.025-1.6 mg.mL\(^{-1}\)), while only the highest concentrations of the essential oils from rainy season leaves (Figures 3S-e and 3S-f) presented anti-inflammatory activity (0.2-1.6 mg.mL\(^{-1}\)). Furthermore, the essential oils showed no toxicity to the transgenic strain of D. melanogaster GMR-GAL4> UAS-Eiger, as verified by the lineage survival test.

A study conducted by Fernandes et al. (2007) also revealed the anti-inflammatory potential of the sesquiterpenes α-bisabolol, β-caryophyllene and α-humulene present in the essential oils in reducing the TNF-α release, in the paw edema of rats. Therefore, since these substances were able to downregulate the cytokine expression and secretion, this finding represents a very promising alternative to be used as a treatment of inflammatory-related diseases.

4. Conclusions

The essential oils from P. myrtoides leaves collected in the dry and rainy seasons presented no significant differences in the quantitative and qualitative profile of volatile constituents and, apparently, did not to affect neither the principal components or the biological activities. In general, the essential oils showed strong to moderate anticycandidial activity for C. glabrata. In the antiprotozoal assay, the seasonal essential oils presented inverted results against two studied parasites. While the essential oils from dry season leaves presented antiprotozoal activities against T. cruzi, the essential oils from the rainy season exhibited antiprotozoal activities against L. amazonensis parasites. Additionally, the essential oils presented antimicrobial efficacy only against S. sanguinis. The sesquiterpenoids present in all the essential oils from P. myrtoides showed a synergistic potential bioactive ability.

In summary, our results supported the antioxidant and anti-inflammatory effects of the essential oils extracted from leaves of P. myrtoides, which vary according to the seasonal period, suggesting that these effects were activated, at least in part, by suppressing the eiger gene, which is orthologous to the human pro-inflammatory cytokine TNF-α.

Thus, P. myrtoides, a plant species from a global biodiversity hotspot (Brazilian Savanna) with potential source of bioactive compounds that offer combined antimicrobial, antiprotozoal, and antioxidant properties, also presents an anti-inflammatory action for further in vivo inflammation models.

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