Research Article

Phytochemical Composition, Antifungal and Antioxidant Activity of Duguetia furfuracea A. St.-Hill

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Background. Duguetia furfuracea is popular plant used in popular medicine. Hypothesis/Purpose. This claim evaluated the phytochemical composition of the hydroethanolic extract (HEDF), fractions of Duguetia furfuracea, and antioxidant and antifungal activity. Methods. The chemical profile was carried out by HPLC-DAD. The total phenolic contents and flavonoid components were determined by Folin-Ciocalteu and aluminium chloride reaction. The antioxidant activity was measured by scavenging of 2,2-diphenyl-1-picrylhydrazyl (DPPH) free radical and ferric reducing ability of plasma (FRAP) methods. The antifungal activity was determined by microdilution assay. Results. HPLC analysis revealed caffeic acid and rutin as major compounds (HEDF), caffeic acid and quercitrin (Mt-OH fraction), and quercitrin and isoquercitrin (Ac-OEt fraction). The highest levels of phenols and total flavonoids were found for Ac-OEt fraction, and the crude extract showed higher in vitro antioxidant potential. The antifungal activity showed synergic effect with fluconazole and EHDF against C. krusei, fluconazole and Mt-OH against C. krusei and C. tropicalis, and Ac-OE and fluconazole against C. albicans. Conclusion. The highest levels of phenols and total flavonoids were marked with antioxidant effect. This is the first report of bioactivity of the synergic effect of HEDF and fractions. More studies would be required to better clarify its mechanism of synergic action.
1. Introduction

The species *Duguetia furfuracea* is a shrub which belongs to the Annonaceae family, being popularly known as “araticum do cerrado,” “ata brava.” The leaves are popularly used for treatment of rheumatism and renal colic, antihiperpipidemic agent, and anorexic agent [1].

Studies with different structures of *D. furfuracea* have pointed out its cytotoxic, bactericidal, and antitumoral properties. Extracts from leaves and roots of *Duguetia furfuracea* presented larvicide activity against *Aedes aegypti* [2]. Alkaloids from peels of subterranean stem showed antitumor, trypanocide [3], and leishmanicidal activities [4, 5]. Extracts from subterranean parts of the plant displayed toxic effect to *Artemia salina* [4]. Our group demonstrated recently the prooxidant and insecticidal activity of the hydroalcoholic extract from leaves of *D. furfuracea* in *Drosophila melanogaster* [6]. Studies regarding antifungal and antioxidant properties of *Duguetia furfuracea* are scarce; thus, the present work contributes to amplifying the knowledge about this species.

The antioxidant potential of the plants has been associated mainly with the presence of phenolic compounds. The chemical structure and reductive properties of these compounds make them active molecules in the scavenging of free radicals and in chelation of transition metals [7]. Secondary metabolites are substances with a generally complex structure playing several roles in the adaptation of plants to the environment; these substances have been attracting interest for their pharmacological and biotechnological properties such as antioxidant [8], anti-inflammatory [9], antimutagenic, anticarcinogenic, gastroprotection [10–12], and antimicrobial [13] ones.

The occurrence of fungal infections is increasing at alarming rates, especially among immunocompromised subjects, such as AIDS patients, transplanted patients, and neonates [14]. Among the pathogens, species of *Candida* are generally associated with these infections, whose incidence is attributed to a variety of factors, either exogenous or endogenous. More than 100 species of *Candida* are known and the frequency of distribution for *Candida* spp. varies in accordance with geographical location [15, 16].

Actually, conventional treatments for fungal infections are not fully effective, since the available drugs lead to secondary effects or to development of resistance [17]. Therefore, the search for new drugs and alternative therapies (including natural products) for the treatment of *Candida* infections has become critical. In this aspect, plants and their derivatives have been contributing to pharmacological research due to their potential as a source for a variety of biologically active ingredients used in drug development. The antimicrobial activity of plants has been identified in some species; however, it should be taken into consideration that besides its beneficial effects, the use of plants may interfere with conventional treatments by interaction with drugs, thus potentiating or minimizing clinical efficacy [18].

Our aim of the present study was to describe the phytochemical characterization, *in vitro* antioxidant potential and to evaluate, for the first time, the antifungal and/or modulatory activity of the hydroalcoholic extract, methanolic and ethyl acetate fractions of *Duguetia furfuracea*.

2. Materials and Methods

2.1. Drugs, Reagents, and Equipment. Sabouraud Dextrose Agar (semisolid) and Sabouraud Dextrose Broth were from Difco Laboratories (Michigan, MI, USA). Dimethyl sulfoxide (DMSO), acetonitrile, and formic, gallic, chlorogenic, ellagic, and caffeic acids were purchased from Merck (Darmstadt, Germany). Antifungal agents fluconazole and nistatina, catchin, quercetin, quercitrin, isoquercitrin, rutin, kaempferol, ethanol, methyl alcohol, hexane, ethyl acetate, Folin-Ciocalteu Reagent, gallic acid, sodium carbonate, aluminium chloride, quercetin, DPPH, ascorbic acid, sodium acetate, TPTZ, ferric chloride, and ferrous sulfate were purchased from Sigma Chemical Co. (St. Louis, MO, USA). High performance liquid chromatography (HPLC-DAD) was performed with the system of HPLC (Shimadzu, Kyoto, Japan), autosampler prominence (SIL-20A), equipped with high-pressure plunger pumps LC-20AT Shimadzu connected to DGU degasser 20A5 with integrator CBM 20A, UV-VIS detector DAD (diode) SPD-M20A, and software solution LC 1,22 SPI. The absorbance measurements were obtained using EnSpire® multimode plate reader (PerkinElmer, USA). All chemical products were of the highest analytical grade.

2.2. Collection of Plant Material. Leaves of *D. furfuracea* were collected from Barreiro Grande, Crato, Ceará (7° 22' 2.8″S, 39° 28' 42.4″W and altitude of 892 m above sea level), Brazil, in September 2011 and identified by Dr. Maria Arlene Pessoa da Silva. A voucher specimen (n. 6703) was deposited in the Herbarium Caririense Dardano de Andrade Lima (HCDAL) of the Regional University of Cariri (URCA).

2.3. Preparation of the Hydroalcoholic Extract, Methanolic and Acetate Fractions. Leaves (1.050 g) of *D. furfuracea* were washed in running water, crushed, and put into glass flasks containing hydroalcoholic solution of extraction (99.8% of ethanol in distilled water) in the proportion of 1:1, for three days. The suspension was filtered, solvent evaporated under reduced pressure, and lyophilized to obtain 261.13 g of crude ethanolic extract. 80 g of this was partitioned with ethyl acetate and methanol to obtain g of 2.28 ethyl acetate fraction (EAF) and 75.6 g of methanolic fraction [19]. The procedure yielded 24.87% for HEDF, 94.5% for methanolic fraction (Mt-OH), and 3.35% for the ethyl acetate (Ac-OEt) fraction.

2.4. Identification and Quantitation of Phenolic Compounds of HEDF by HPLC. Mt-OH and Ac-OEt fractions were submitted to the chromatographic analysis of reversal phase. The chemical composition of the HEDF was previously determined by our group [6] using the same procedure described in this section.

The chromatographic analyses were performed under the same gradient conditions using the column C18 (4.6 mm × 250 mm) charged with particles of 5μm of diameter. The mobile phase was water containing 1% of formic acid (A), acetonitrile (B) and the composition gradient was 13% of B...
fluoride 

3.5. Determination of Total Phenols. The quantification of phenolic compounds was performed using the Folin-Ciocalteu method that involves the reduction of the reagent by phenolic compounds present in the samples forming a blue complex whose intensity increases linearly at 760 nm, as described by Swain and Hillis (1959) [21]. For the assays, 4 µL of samples (HEDF, Mt-OH, and Ac-OEt) at a concentration of 103 µg/mL to an incubation medium consisting of 0.1N Folin-Ciocalteu reagent and 1.25% (w/v) Na2CO3 in a final volume of 284 µL. After two hours of incubation in the dark, at room temperature, the absorbance was measured at 760 nm. The experiments were carried out in triplicate. The index of total phenolic compounds was expressed as equivalents of gallic acid per gram of the sample (mg GAEq/g), calculated through a curve of gallic acid, built with concentrations ranging from 50 to 500 µg/mL.

2.6. Determination of Total Flavonoids. The quantification of flavonoids was made according to Quettier-Deleu et al. 2000 [22]. The method is based on the measurement of absorbance, at 415 nm, of the complex formed between flavonoid compounds and aluminum cation in ethanol. For the assays, samples were incubated with AlCl3 (2%) in a 1:1 reaction. The final volume of the reaction medium was 300 µL. The experiments were carried out in triplicate. The index of total flavonoids was expressed as equivalents of quercetin per gram of the sample (mg QEq/g), calculated through a curve of quercetin, built with concentrations ranging from 0.625 to 25 µg/mL.

2.7. Antioxidant Activity

2.7.1. Scavenging Activity of the DPPH Radical. The antioxidant activity of the extract and fractions was also checked by the DPPH method as described by Brand-Williams et al. (1995) elsewhere [23] with minor changes. This test is based on the reduction of the stable free radical DPPH which presents a deep violet color in solution and turns to a yellowish color when neutralized. The mixture of the reaction was composed of 50 µL of samples (extract and fractions), 50 µL of solvent, and 100 µL of solution and 0.3 mM of the radical DPPH in ethanol. The measurement of the absorbance was at 517 nm after 30 minutes. The samples (HEDF, Mt-OH, and Ac-OEt) were diluted in ethanol and water (1:1) and the standard substance, ascorbic acid, was diluted in water. All tests were made in triplicate. Results were expressed as IC50 values defined as the concentration of antioxidant required to sequester 50% of the DPPH radicals. IC50 was calculated by nonlinear regression.

2.7.2. FRAP (Ferric Reducing Antioxidant Power). The FRAP assay was conducted as described previously by Benzie and Strain (1996) [24]. FRAP solution consisted of 10 mM TPTZ, 20 mM ferric chloride in acetate buffer 0.3. In 96-well microtiter plate, we added 9 µL of the samples (HEDF, Mt-OH, and Ac-OEt); 27 µL of water; and 270 µL of the FRAP solution. After incubation at 37 °C for 30 minutes absorbance was taken at 595 nm. Samples readings were compared to a ferrous sulfate II standard curve. Results were expressed as µM ferrous sulfate II (FeSO4) equivalents per gram of sample. Experiments were done in triplicate.

2.8. Antifungal Activity

2.8.1. Culture Medium and Inoculums. The antifungal activity was evaluated using the standard fungal strains C. albicans (ATCC 40277), C. krusei (ATCC 6438), and C. tropicalis (ATCC 40042) donated by the Universidade Estadual da Paraíba. In the biological tests, we used the following culture medium: Sabouraud Dextrose Agar (semisolid) and Sabouraud Dextrose Broth (liquid) prepared according to the manufacturer’s specifications. Fungal cultures kept at 4°C were transported to the Sabouraud Dextrose Agar medium and incubated at 35°C for 24 hours. As to the preparation of the inoculum, the pricked out strains were transferred to the sterile saline solution (0.9% NaCl), composing of a fungal suspension (inoculum) until obtaining the concentration of 105 UFC/mL according to the scale of McFarland [25].

2.8.2. Minimum Inhibitory Concentration Test: CIM and the Modulation of Standard Antifungal Action. The method of microdilution in sauce was used to determine the minimum inhibitory concentration (CIM). The samples (HEDF, Mt-OH, and Ac-OEt) were weighed and solubilized initially in
dimethyl sulfoxide (DMSO) and diluted at 1024 μg/mL using sterile distilled water (test solution).

We distributed 100 μL of inocula, prepared previously, in each cavity of a 96-well microtiter plate and, thereafter, it was submitted to a serial double dilution using 100 μL of the samples with concentrations that range from 1024 to 0.5 μg/mL. The plates were transported to the incubator for 24 hours at 35°C [26]. The identification of CIM was performed through the visual observation of the turbidity provoked by the fungal growth, with the CIM being defined as the lowest concentration of the sample in which no fungal growth was observed [25].

To observe how these samples could affect the action of the standard antifungal agents against the strains tested, we used the method proposed by [27]. The extract and the fractions were tested using a subinhibitory concentration (MIC/8 = 64 μg/mL). We distributed, in each well, 100 μL of solution containing 1.675 μL of liquid medium (Sabouraud Dextrose Broth) 10%; 200 μL of inoculum (fungal suspension); and 125 μL of the natural product (extract and fractions). After that, 100 μL of the antifungal agents was added to the first cavity and following the serial dilution along the other cavities. The concentrations of the antifungal agents ranged from 1024 to 0.5 μg/mL. The tests were performed in triplicate.

### Statistical Analysis

The results of the tests were done in triplicate and expressed as geometric mean [28]. Statistical differences between samples were tested by analysis of variance ANOVA followed by Tukey’s or Dunnett’s post hoc test when necessary. The differences were considered significant when P < 0.05.

### Results and Discussion

#### 3.1. Identification and Quantification of Phenolic Compounds by HPLC.

The chromatographic and spectral profile of Mt-OH and Ac-OEt fractions revealed the presence of gallic acid (t_R = 9.95 min; peak 1), catechin (t_R = 16.08 min; peak 2), chlorogenic acid (t_R = 20.14 min; peak 3), caffeic acid (t_R = 24.63 min; peak 4), ellagic acid (t_R = 37.29 min; peak 5); rutin (t_R = 39.87 min; peak 6); isoquercitrin (t_R = 44.93 min; peak 7); quercitrin (t_R = 48.15 min; peak 8); quercetin (t_R = 51.07 min; peak 9); and kaempferol (t_R = 61.56 min; peak 10).

The main compounds present in the Mt-OH fraction were caffeic acid (32.47 ± 0.03 mg/g) and quercitrin (31.56 ± 0.01 mg/g) while gallic acid (5.29 ± 0.01 mg/g) and isoquercitrin (5.49 ± 0.02 mg/g) were the least abundant [6].

The determination of total phenols and flavonoids was higher in Ac-OEt fraction (657.05 mg/EAG/g and 120.9 mg EQ/g, resp.) while gallic acid (20.05 ± 0.01 mg/g) and isoquercitrin (18.73 ± 0.01 mg/g) were the least present (Figure 1 and Table 1). The chromatographic profile of HEDF demonstrated the presence of caffeic acid and rutin as major compounds (31.96 ± 0.03 mg/g and 120.9 mg EQ/g, resp.) while gallic acid (5.29 ± 0.01 mg/g) and catechin (3.16 ± 0.02 mg/g) were the least abundant [6].

The determination of total phenols and flavonoids was shown in Table 2. The content of total phenols and flavonoids was higher in Ac-OEt fraction (657.05 mg/EAG/g and 120.9 mg EQ/g, resp.), followed by Mt-OH fraction (289.33 mg/EAG/g and 76.26 mg EQ/g, resp.) and HEDF (231.26 mg EAG/g and 87.57 mg EQ/g, resp.). It is recognized that flavonoids are preferably extracted by ethyl acetate solvent.

The *in vitro* antioxidant potential of crude extract (HEDF) and fractions of *D. furfuracea* was evaluated by two different methods, FRAP, which measures the ferric reducing antioxidant power of compounds, and ability of sequestering
the synthetic radical DPPH. The crude extract (HEDF) presented the best ferric reducing potential (166.73 ± 5.13 μM of Fe^{2+}/g of sample), followed by Mt-OH (126.43 ± 4.98 μM of Fe^{2+}/g of sample) and Ac-OEt fractions (118.20 ± 1.08 μM of Fe^{2+}/g of sample) (Table 3). The potential of scavenging of radical DPPH was expressed as IC_{50} in μg/mL of extract or fractions and compared with the positive control ascorbic acid. HEDF presented the better antioxidant activity in the DPPH test with IC_{50} values of 33.15 μg/mL when compared to Ac-OEt (39.32 μg/mL) and Mt-OH (42.32 μg/mL).

In this study, the antifungal potential of the hydroalcoholic extract of *D. furfuracea* (HEDF) and methanolic (Mt-OH) and ethyl acetate (Ac-OEt) fractions was tested against standard strains of *C. albicans*, *C. tropicalis*, and *C. krusei*. According to the results, both extract and fractions presented minimal inhibitory concentration (CIM) ≥1024 μg/mL against all the fungi strains tested. However, the extract and the fractions of *D. furfuracea* presented synergic effect when they were associated with fluconazole, indicating its modulatory action against fungi when associated with clinically relevant drugs. The HEDF and Mt-OH fraction potentialized the effect of the fluconazole when tested against *C. krusei* as observed in Figures 2(a) and 2(b). The methanolic fraction also presented synergism with fluconazole against *C. tropicalis* (Figure 2(b)) and Ac-OEt fraction potentialized the effect of fluconazole against *C. albicans* (Figure 2(c)).

4. Discussion

Phenolic compounds and some of their derivatives are known by their antioxidant properties. The antioxidant activity of some medicinal plants is correlated to the total phenolic and flavonoids indexes [29]. The level of total phenols for *D. furfuracea* extract and fractions is comparable with
other Brazilian medicinal plants of *Duguetia* genus [30, 31]. Although the total index of phenols and flavonoids has been higher in the fractions than in crude extract, it is possible to connect these compounds with *in vitro* antioxidant activity, as determined by FRAP and DPPH methods. Though the *in vitro* antioxidant activity was higher in the crude extract than the fractions, this analysis suggests that other compounds in the crude extract of *D. furfuracea* contribute to its more effective antioxidant activity [32]. In a previously published study by our group, the crude extract (HEDF) was demonstrated to present alkaloids in its phytochemical constitution [6]. In this work we can see a direct correlation between the concentration of flavonoids and antioxidant FRAP activity ($r = -0.801$). Those results were confirmed with others results present in literature [33–35].

The primary mechanism of fluconazole’s action occurs by the inhibition of the fungal enzyme lanosterol 14α-demethylase (CYP51), which is a cytochrome enzyme P-450, involved in the synthesis of the ergosterol, the most important sterol in the fungal cell membrane [36]. It is known that many medicinal plants may modulate the activity of several antimicrobial agents [37, 38]. In a previous study, the aqueous extract of

![Figure 2](image-url)

**Figure 2:** Modulatory activity of (a) HEDF (hydroalcoholic extract of *D. furfuracea*), Mt-OH, and Ac-OET against the fungi *C. albicans*, *C. krusei*, and *C. tropicalis* (concentrations ranging from 1024 μg/mL to 0.5 μg/mL). $P < 0.001$ related to control group; (b) Mt-OH (methanolic extract of *D. furfuracea*) against the fungi *C. albicans*, *C. krusei*, and *C. tropicalis* (concentrations ranging from 1024 μg/mL to 0.5 μg/mL); (c) Ac-OEt (ethyl acetate extract of *D. furfuracea*) against the fungi *C. albicans*, *C. krusei*, and *C. tropicalis* (concentrations ranging from 1024 μg/mL to 0.5 μg/mL). Statistical analysis: one-way ANOVA followed by Student-Newman-Keuls test. **$P < 0.001$** versus fluconazole.
Table 3: Antioxidant activity of HEDF and fractions of D. furfuracea.

| Samples     | Sequestering of the radical DPPH EC50 (µg/mL) | FRAP µM Fe2+/g of the sample |
|-------------|---------------------------------------------|-----------------------------|
| HEDF        | 33.15b                                      | 166.73 ± 5.13a              |
| Mt-OH       | 42.32c                                      | 126.43 ± 4.98b              |
| Ac-OEt      | 39.32c                                      | 118.20 ± 1.08b              |
| Asc. Ac.    | 17.50a                                      | —                           |

The values were expressed as mean ± SD (n = 3); EFe2+ = equivalent of iron; HEDF (hydroalcoholic extract of D. furfuracea); Mt-OH (methanolic fraction); Ac-OEt (ethyl acetate fraction); and Asc. Ac. (ascorbic acid). Results are expressed as mean ± SEM (n = 3). Averages followed by different letters differ by Tukey’s test at P < 0.05.

the leaves and fractions of D. furfuracea when combined with aminoglycosides presented synergistic activity against Escherichia coli and Staphylococcus aureus [39].

Phenolic compounds and flavonoids have demonstrated potential therapeutic activities as antifungal, antibacterial, and antioxidant agents [40, 41]. Although the mechanisms underlying antimicrobial pharmacology of the phenolic compounds are rather variable, many of them act by promoting damage to the function of the cell membrane or cell wall [42]. The analysis by HPLC of the extract and fractions of D. furfuracea, as described previously, revealed the predominance of the following compounds: caffeic acid, chlorogenic acid, rutin, quercitrin, and isoquercitrin. There is a study that demonstrated that chlorogenic acid presented antifungal activity against the yeast of the gender Candida [43]. Six flavonoids that were isolated from plants, among them, rutin, presented antibacterial and antifungal activity [40].

Sun et al. (2004) showed the influence of phenolic compounds in fluconazole antifungal properties. This paper shows that the concentration of fluconazole in C. albicans was found to be increased with the increment of the phenolic compounds concentration when they were in combination. This result corroborated synergetic activity present in this work [44].

It is possible to speculate that some of these chemical constituents, especially the flavonoids, are responsible for the pharmacological properties found. However, the isolation and the activity of alkaloids and acetogenines have stood out in studies with plants of the Annonaceae family. The biological activity as antimicrobial capacity and antioxidant activity present in A. muricata can be attributed to the presence of acetogenines [37, 45]. Alkaloids as aminophenols present the bark of Annona salmammatii D. C. was responsible for the antioxidant and antimicrobial capacity [46].

5. Conclusion

We can conclude that the crude extract of D. furfuracea (HEDF) and its methanolic (Mt-OH) and ethyl acetate (Ac-OEt) fractions have an important antioxidant activity (in vitro) when compared to other natural compounds. The crude extract (HEDF) presented highest antioxidant activity in vitro when compared to Mt-OH and Ac-OEt, as determined by DPPH and FRAP methods. However, it was not possible to observe a positive correlation between the antioxidant activity and the total index of phenols and flavonoids identified, indicating that compounds other than phenolics may contribute to the antioxidant potential of the plant extracts. The crude extract and fractions of D. furfuracea presented a synergistic activity with fluconazole when tested against strains of C. albicans, C. tropicalis, and C. krusei, indicating a potential antifungal activity via modulation of clinically used drugs against fungal infections. More studies are needed to clarify the mechanisms involved in this phenomenon as well as other potential biomedical and biotechnological applications of D. furfuracea.

Abbreviations

HEDF: Hydroethanolic extract of Duguetia furfuracea
Mt-OH: Methanolic fractions of Duguetia furfuracea
Ac-OEt: Acetate fractions of Duguetia furfuracea
HPLC DAD: High performance liquid chromatography with diode array
DPPH: 2,2-Diphenyl-1-picrylhydrazyl
FRAP: Ferric reducing antioxidant power

Competing Interests

All authors wish to confirm that there is no known conflict of interest associated with this publication and there has been no significant financial support for this work that could have influenced its outcome.

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