Evaluation of Antigonorrhea Activity and Cytotoxicity of Helichrysum caespititium (DC) Harv. Whole Plant Extracts

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Abstract

Over 80% of African population depends on traditional knowledge for their well-being, and especially on plants as medicines. Although Helichrysum caespititium is among plants that are commonly used by traditional healers in Africa, its biological activities are still not scientifically proven and reported. The primary objective of this study is to assess the antigonorrhea activity and cytotoxicity of H. caespititium whole plant. The plant material was subjected to a serial exhaustive extraction to obtain different solvent extracts using n-hexane, dichloromethane, methanol, and water. The antigonorrhea activity of the four plant extracts (n-hexane, dichloromethane, methanol, and water extracts) against 2008 WHO Neisseria gonorrhoea reference strains and the toxicity of the extracts against rat liver cells were investigated. All four H. caespititium extracts showed good activity against the four 2008 WHO N. gonorrhoea strains (F, O, N, G strains) under study in the range of 0.037 to 0.33 mg/ml. n-Hexane extract was observed to be the most potent against all the four strains with a lowest Minimum Inhibitory Concentration (MIC) value of 0.037 ± 0.0 mg/ml against G strain, which was comparable to gentamicin (standard 1) and more active compared to amoxicillin (standard 2), and also the most less toxic of all with LC₅₀ value of 428.77 ± 4.76 μg/ml followed by water extract (394.36 ± 5.41 μg/ml) and methanol (357 ± 2.81 μg/ml). The results justify the usage of H. caespititium in the traditional medicine against gonorrhea infections.

Keywords: Antigonorrhea; Cytotoxicity; H. caespititium; MIC

Introduction

Over 80% of African population depends on traditional knowledge for their well-being, and especially on plants as medicines. The Southern African region contains more than 350,000 species of flowering plants, in which many of them have been used by traditional healers. However, their biological activities are still not scientifically proven and reported. Among the 500 species that contain Helichrysum family, which are discarded worldwide, considerable species, approximately 245-246, occurs in Africa and Madagascar [1-4]. Helichrysum caespititium is one of those plants growing in Southern African region where it is referred to as inpepe (Zulu), impelho (isiXhosa), seledsa-phoko (South Sotho), moriri-wa-naha (Kvena), and sepahaney (Kgatla).

Helichrysum species name “caespititium” was derived from the Latin word “caespitose,” which means very much tufted, matted, referring to the cushion-forming growth habit. The plant has been referred to as everlasting [1]. Botanically, H. caespititium (DC) is presented as a prostate, perennial, mat-forming herb that is profusely compact shrublet with branched and densely tufted. Leaves are scattered with orange silvers. Silvery white flowers appear in late summer with yellow centers and pale furry underneath [1].

The plant has been used since ancient time for treatment of several diseases, such as bronco-pneumonia, tuberculosis, and intestinal ulceration. Moreover, it is used in styptic wound dressing particularly during the circumcision rites [5], bruises, cuts, and sores [3]. Furthermore, the plant has been involved in the treatment of skin infection diseases, respiratory problems, gastro-intestinal tracts, and diarrhea in Sekhukhune and Waterberg municipality districts in the Limpopo province, South Africa [6]. The Basotho population inhales the smoke emerging from burning of H. caespititium plant material for the treatment of headache, chest colds as well as for the treatment of internal wounds such as intestinal ulceration. Moreover, the concoction of H. caespititium has been drank by Bakwena and Bakgatla populations in ancient time for the treatment of gonorrhoea infection [1].

Gonorrhea is a common sexually transmitted disease that affects thousands of men and women annually, particularly in the United States [7]. Although gonorrhea is easily treated, it can cause serious and sometimes enduring complications such as pelvic inflammatory disease in women and epididymitis and barrenness in men [7-9]. Regimens for the treatment of gonorrhea are increasingly being based on oral and/ or injectable expanded-spectrum third generation cephalosporins such as cefixime and ceftriaxone, but worries have recently been uttered about their continuing efficacy [10-15]. This condition, as well as the emergence of reduced susceptibility and resistance to azithromycin, has called for improved efforts for the control of gonococcal disease [12,14,15].

In our endeavor to find cure for infectious disease particularly gonorrhea, we decided to investigate the claimed antigonorrhea activity of H. caespititium by traditional healers and the cytotoxicity of H. caespititium plant extracts. Although the antibacterial activity of the plant has been proven by Mathenge et al. [5], a search in the literature reveals that H. caespititium plant’s antigonorrhea activities and cytotoxicity have not yet been studied and proven scientifically. As there has been a concern about the efficacy of some current antgonorrhea drugs toward gonorrhea infections [10-15], it is of considerable urgency to find other drugs that can surmount the difficulty experienced at present.
In this study, to the best of our knowledge, we are the first to report the antigonorrhea activities of the plant against 2008 WHO *Neisseria gonorrhoea* reference strains and the cytoxicity of the plant extracts.

**Materials and Methods**

**Material**

The solvents that were obtained from Sigma (South Africa) for extraction were \( n \)-hexane, dichloromethane, and methanol (reagent grade). The water was purified from water distillation plants in our laboratory. All other chemicals were of analytical grade or GC grade.

**Collection and identification of the plant**

The whole plant material of *Helichrysum caespititium* was collected from Masealama village, which is situated at 29.88° East longitude and -23.83° South latitude in Polokwane Municipality, Capricorn District Municipality in the Limpopo Province, South Africa. The plant was then taken to the South African National Biodiversity (SANBI) in Pretoria for identification, and the identification code is DTH 9006000.

**Processing of plant material**

The collected plant material was dried at room temperature before being grinded into powder using Mellerware Coffee Bean Grinder machine (Aromatic, 29105A, South Africa). The resulted powder was kept in dark at 4°C for further usage.

**Extraction**

A mass of 160 g of powdered plant material was subjected to a serial exhaustive extraction using the maceration method in 3000 ml of \( n \)-hexane, dichloromethane, methanol, and water (starting with less polar to more polar solvent). The mixtures were shaken for 24 h at 120 rpm using Labtech shaker. The filtration was performed using the Whitman filter paper No. 1. The extracts were concentrated using rotary evaporator with reduced pressure at temperature up to 40°C. Each solvent used in the extraction was repeated three times, and the extracts were combined. Moreover, the extracts were used in this study.

**Microorganisms**

The MIC test involved four *N. gonorrhoea* strains that are used by WHO as reference strain for global quality assurance and quality control of gonococcal antimicrobial resistance testing, and the strains are identified as F, N, O, and G [16]. The selection of the 2008 WHO *N. gonorrhoea* strains in this study was based on their type. 2008 WHO *N. gonorrhoea* strains have eight strains among which two are African type (M and O strains), one is an Asian type (N strain), one is Dutch type (G strain), three are wild-type (F, L, and P strains), and one is an unknown type (K strain). It was decided to use one strain in each type (G strain), three are wild-type (F, L, and P strains), and one is an Asian type (N strain), one is Dutch type (M and O strains), one is an African type (A strain), one is an unknown type (K strain). It was decided to use one strain in each type namely F (wild-type), G (Dutch type), N (Asian type), and O (African type). The adopted experimental procedure was that proposed by Eloff [17] and that from the Clinical and Laboratory Standards Institute (CLSI) [18], but with slight modification. Hundred and fifty microliter (150 μl) of Mueller Hinton broth was pipetted into each well of the 96-wells plate. Thereafter, 75 μl of the extract solution (\( n \)-hexane, dichloromethane, methanol, or water) (1 mg/ml) was prepared in water and 0.5 ml of acetonitrile to allow the extracts to dissolve, because it does not completely dissolve in water alone. This was added to first wells followed by a threefold serial dilution. Thirty five microliters (35 μl) of an overnight bacterial suspension grown on New York City plate (GC-agar) was added to each well. The microtiter plate was sealed and then anaerobically incubated for 24 h at 37°C in the presence of 5-10% of CO₂ for the survival of the bacteria under study. After incubation, 40 μl of p-jodinitrotetrazolium chloride (INTC) (0.1 mg/ml) was added to each well, and then the plate was further incubated for approximately 40 min to 1 h at the similar condition. Gentamicin and amoxicillin solutions were used as positive control. The wells that showed pink color explained the growth of bacteria. However, the yellow colored wells or colorless wells explained no bacteria growth (formation of formazan). The MIC was recorded as the lowest concentration that could not produce the visible bacteria growth.

**Cytotoxicity assay**

The extracts of *H. caespititium* were screened for cytotoxic activity in H411E rat hepatoma (liver) cell lines using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) assay [19,20]. Cells were maintained in minimum essential medium (MEM) supplemented with 10% fetal calf serum (FCS) and 1% penicillin/streptomycin solution. Previously established 80% confluent monolayer was trypsinized and resuspended in fresh MEM for seeding at a density of 1.0 × 10⁵ cells/ml into each well (100 μl) of 96-well plate for the toxicity assays. Following overnight incubation at 37°C in a 5% CO₂, 100 μl of fresh media containing plant extracts was added to the cells in the plates for cytotoxicity assay. Prior to addition of fresh medium with plant extracts, stock solutions (100 mg/ml) were prepared and further serial dilutions with growth media to six concentrations were made and used for the assays. DMSO was used as solvent (negative) control, while doxorubicin was used as the positive control. The cells were then incubated for 24 h. After the incubation, the medium was removed by aspiration and fresh media together with 20 μl of MTT (5 mg/ml in phosphate buffered saline, PBS) were added into each well. After a further incubation period (4 h), the medium was carefully aspirated without disturbing the MTT crystals at the bottom of the wells and replaced with 40 μl of undiluted DMSO. The concentration of MTT reduced was measured at 570 nm after gentle shaking. The wells containing only medium and MTT was used to blank the microplate reader (Epoch BioTek). The LC₅₀ calculated from a linear curve of log of concentrations versus average absorbances represents the concentration of extract that resulted in a 50% reduction of absorbance in comparison to the untreated cells.

**Statistical analysis**

The results of all the experiments involved in this study were performed in triplicate. One-way analysis of variance (ANOVA) followed by the t-test were used in the data analysis. Therefore, all results are presented as mean values ± standard deviation (SD). All P values lower than 0.05 were considered as significant (p < 0.05).

**Results and Discussion**

All four plant extracts tested were active against the four WHO *N. gonorrhoea* strains with MIC values ranging from 0.037 to 0.3 mg/ml (Table 1). It is stipulated that the antimicrobial activity of a crude extract is considered significant when the MIC is below 100 μg/ml (0.1 mg/ml), moderate when between 100 and 625 μg/ml (0.1 and 0.625 mg/ml), and low when more than 625 μg/ml (0.625 mg/ml) [21,22]. For pure compounds, the activity is considered significant when the MIC is below 10 μg/ml (0.01 mg/ml), moderate when between 10 and 100 μg/ml (0.01 < MIC < 0.1 mg/ml), or low when greater than 100 μg/ml (0.1 mg/ml) [21,22]. Moreover, according to Gibbons *et al.* [23], the values of MIC below 1 mg/ml for extracts and 64 μg/ml (0.064 mg/ml) for single chemical entities are considered significant. Therefore, these results (Table 1) are worth considering.
Plant extracts/standards & MIC (mg/ml) & Cytotoxicity (μg/ml) \\
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<th>F</th>
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<tr>
<td>H.E</td>
<td>0.33 ± 0.00</td>
<td>0.037 ± 0.00</td>
<td>0.33 ± 0.00</td>
<td>0.108 ± 0.067</td>
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<td>D.E</td>
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<td>M.E</td>
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<td>W.E</td>
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<td>Amoxicillin truth</td>
<td>&gt;0.33</td>
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<td>Gentamicin &amp; hexane extract</td>
<td>0.22 ± 0.11</td>
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<td>0.22 ± 0.11</td>
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<tr>
<td>Doxorubicin</td>
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In conclusion, this study managed to reveal some truth about the plant antigonorrhea activities and toxicity character toward cells. The plant in general is a strong antigonorrhea agent with n-hexane extract being the most compared to other extracts (dichloromethane, methanol, and water extracts). In addition, the plant can be used to address the problem of gonorrhea infections like other plant extracts that have been published elsewhere [24]. Looking at the cytotoxicity character of the plant extracts, this plant is proven to be nontoxic to cells. 

This study is a preliminary study toward the isolation of the phytochemical compounds responsible for the claimed bioactivity of *H. caespititium* plant. An in-depth study of the extract in terms of structure elucidation of the bioactive compounds is under way to provide a good base for all the phytochemical functions mentioned above and their bioactivity studies.

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References


