Research Article

Antibacterial and antifungal activities of extracts of Convolvulus leiocalycinus and Haloxylon griffithii of Balochistan, Pakistan

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Abstract
Different medicinal plants have the potential of antibacterial and antifungal activities. Therefore, this research was executed to assess the antibacterial and antifungal activities of different fractions i.e. water (H₂O), chloroform (CHCl₃), n-hexane (C₆H₁₄) and ethyl acetate (C₄H₈O₂) of Convolvulus leiocalycinus and Haloxylon griffithii by applying Agar well diffusion method. Bacterial strains of four types i.e., 3 gram-negative and 1 gram-positive namely Escherichia coli, Klebsiella pneumonia, Staphylococcus aureus, Pseudomonas aeruginosa and a fungal strain like PBF-1 were used to investigate the antibacterial and antifungal activities in the selected plant extracts. The water, chloroform, ethyl acetate and n-hexane fractions of C. leiocalycinus exhibited antibacterial activities of 7, 7, 7 and 6mm zone of inhibition respectively against S. aureus bacteria. Consequently, the same four fractions of H. griffithii displayed antibacterial activities against S. aureus 8, 8, 9 and zero mm zone of inhibition respectively. Considering the antifungal activity, the ethyl acetate fraction of C. leiocalycinus showed strong inhibition i.e., ++. But in H. griffithii, only the n-hexane fraction gave partial inhibition in antifungal activity. These results indicated that the ethyl acetate fraction of C. leiocalycinus exhibited strong antifungal activity as compared with the same fraction of H. griffithii showing that C. leiocalycinus is more potent as compared to H. griffithii. This justifies the usage of both plants in traditional medicines in future. It is also suggested that the use of entire extract of both the plants in foods can improve their shelf life.

Keywords: Agar well diffusion; Antibacterial activity; Antifungal activity; fungal strain PBF-1.

Introduction
Although, immense development in human medication, infective disorders produced by viruses, fungi, parasites and bacteria are yet a major risk to the health of humans. The effects are far more savior on developing nations by reason of the absence in development of widely spread drug resistance [1]. The resistance of drugs over the last 2 decades is in development along with the undesirable side effects. Appearance of some antibiotics [2] has directed to the discoveries of the modern antimicrobial agents in plant extracts with
the purpose of discovering new chemical structures, converting above drawbacks [3]. World’s 80% of available medicines utilized in developing countries are from medicinal plants [4]. A wide-ranging variety of medicinal plants utilized in the extraction as raw drugs having different medicinal properties. Local communities collect very minimum amounts of the raw drugs and used for local use by local healers, various other drugs are collected in huge scale and merchandised in market, which for many herbal industries are raw material [5]. Clinical microbiologists are very interested in screening of medical medicine for new treatments [6]. The active principle of numerous drugs originate in the plants are secondary metabolites. The antimicrobial activities from extracts of plants are existing in different components, phenolic compounds and aldehydes are also included [7]. Drug resistance preparation against normally used antibiotic in human pathogens needs to find latest antimicrobial substances including plants from other sources [8].

Resistance of bacteria against antibiotics has become an increasingly global concern [9]. Extreme caution doctors believe that bacteria which are resistant towards antibiotic an important or major issue in treating patients [10]. Resistance of bacteria is encouraging to resume in the role of antimicrobial research herbs against resistant strains [11, 12]. A large quantity of medicinal plants has been known as a beneficial resource for natural antimicrobial compounds [13]. Medicinal plant extracts provide sufficiently potential for growth of new agents which is effective against contaminations formerly problematic to treat [14].

A large extent antibiotics utility is the development of microbe resistance and this resistance is spreading around the globe [15, 16]. The result of the failure of the resulting therapy increases. Another limit for the use of antibiotics is the rising price [17, 18]. Synthetic organic compounds are more than 99% but still natural products are more than the third part of all drug sales [19]. The high means of antimicrobial molecules are the availability of medicinal plants. A number of medicinal plants extracts are used to treat many diseases because they have potential protection activity. Some of these bioactive molecules are screened and traded in a market like raw substance for many herbal industries [20]. After observing more side effects of artificial drugs compared to their benefits, experts focused on gaining benefits from medicinal plants [21]. Out of 422127 plant species have been reported worldwide, it is estimated that approximately 35000 to 70000 plant species are used as medicinal plants [22]. Approximately, 80% population of rural areas in Pakistan depends on traditional medicines [23].

Medicinal plants can provide abundance of antimicrobial agents and has been examined for hundreds of biological activities. Small amounts of crude materials are collected by local people and they use them to cure infections. Raw material is also collected on a large scale and is sent to deliver trade in the market to the herbal industries [5]. Parasites and pathogens will be the biggest hazard to humans. Pathogens are allowed to be established in new areas due to climate change. Pathogens and their vectors are developing many resistances made compounds used to manage them. The rising incidence of multidrug resistant strains of bacteria face challenges for treating recent emerging bacterial infections with less sensitivity for antibiotics [24]. In present research, the two medicinal plants like H. griffithii and C. leiocalycinus were investigated for their antimicrobial and antifungal activities so that their therapeutic natures are disclosed to the researchers.

H. griffithii belongs to the family of Chenopodiaceae, familiar with name of Cat tail Family. It covers 100 genera and 1200 species [22]. The class of Haloxylon covers 13 species growing in dry places of the North-African and Arabian deserts and
South-West Asia. The species of the genus Haloxyton are shrubbery [23]. Five species of this class are found in Pakistan [24]. Consequently, *C. leiocalycinus* belongs to the family of Convolvolaceae. However, *Convolvulus*, a genus, comprising of 200-250 species of flowering plants. *C. leiocalycinus* is found in Balochistan on stony slopes at the height of 1500m, also found in the hilly areas of Hanna Lake, Ziarat, Brewery close to Quetta city.

**Materials and methods**

**Plant materials and sample collection**

From Hanna valley, situated in the north of Quetta and Spini road plants part of *C. leiocalycinus* and *H. griffithii* specimens were collected on May 9, 2017. The specimens were recognized by “Prof. Dr. Rasool Bakhsh Tareen”, Taxonomist, University of Balochistan. Both plants were dirty with dust and thus cleaned by tap water, to save them from bacterial and fungal attack and they were air dried for half month. To maintain their metabolites, they were protected from sun rays. They were subjected to determine antifungal and antibacterial activities as long as they were dried.

**Instruments and reagents**

Grinder, rotary evaporator, separating funnel, beakers, round bottom flask, conical flasks, petri dishes, auto clave, laminar flow cabinet, cotton swab, wire loop, burner or spirit lamp, aluminum foil, electronic balance, refrigerator, Cork borer, tweezers, ethanol, methanol, n-hexane, ethyl acetate, chloroform, MHA (Mueller Hinton Agar) and distilled water were utilized in present research.

**Extracting solvent**

Solvent extraction is a method in which plants are dissolved in specific solvent to extract vital medicinal ingredients from plants. The roots, leaves and stems of the *C. leiocalycinus* and *H. griffithii* were air dried and grounded to fine powder. Separated the thick part from the powder ones and was poured to grinder for grinding. The fine powder of 5kg soaked in 15L of methanol for a week with shaking and infrequent stirring. To filter the mixture, the filter paper (Whatman No.1) was set. At least 3 times the extraction was done until no color changed noticed. The filtrate was evaporated on rotary evaporator. The pressure was reduced at 35°C to obtain a partly dehydrated crude methanolic extract (CME). Weighed a china dish and then the methanolic crude extract was poured into it. The crude was allowed for evaporation in cold, arid and dark place. The residue weighted 300g after the evaporation of the residue. This CME was used for investigation of antifungal and antibacterial activities of *C. leiocalycinus* and *H. griffithii*.

**CME fractionation**

A process of classification of analysis wherein the dissolved compounds of CME are kept aside into various parts based on their translation & polarities. For this, the best choice is the solvent extraction wherein the compounds are fractioned in to two parts not forming similar mixture based on their dissolvability. Usually in CME classification of analysis, the solvent selected are water (H₂O), chloroform (CHCl₃), n-hexane (C₆H₁₄) and ethyl acetate (C₄H₇O₂) relating to reduction in their polarity. The dissolved material utilized were of analytical grade. In the initial stage of segregation, the compound was segregated into non-polar, moderate polar and extremely polar. Thus, for classification of analysis, 60g of CME is grinded into powder than mixed to distilled water of 200mL and poured into separating funnel. Then poured 400mL of n-hexane into funnel, closed with lid and shook for 15-20 minutes. To release the pressure during shaking course, the separating pipe was released occasionally. Kept the separating funnel for fifteen minutes to segregate the two layers after shaking. Collected n-hexane layer in flask. The process was done again 3 times (3x400 mL) to ensure no another compound going into layer of n-hexane to obtain the crude fraction of n-hexane. Using vacuum rotary evaporator (RV06-ML, IKA WERKE,
GERMANY), the n-hexane was evaporated at 100 rpm and below 15°C temperature. Weighed the crude and kept in freezer at -20°C in a labeled flask.

From earlier step, water fraction taken in separating funnel and chloroform (400mL) was used into the funnel. For 15-20 minutes the two solvent were mixed, occasionally released the separating funnel during this mixing. After the process, the funnel was kept still for 10 to 15 minutes so that two layers are entirely formed. In a separate flask, the chloroform layer was collected and two more times repeated the same step to complete extraction of solvent so that no more compounds get into water solution of chloroform layer. To get the chloroform fraction (3x400mL) a total of 1200mL of chloroform was used. At 100 rpm and below 25°C temperature, the fraction of chloroform was evaporated on vacuum rotary evaporator. The crude fraction of chloroform was weighed and kept in freezer at -20°C in a labeled flask.

With water fraction, the analytical method proceeds by mixing it with ethyl acetate. First of all, in separating funnel 400mL of ethyl acetate was added to water layer and were mixed for 20 to 25 minutes. Occasionally released the separating funnel to release inside pressure of funnel during shaking. For about 15 minutes, the funnel was kept still so the two immiscible layers could be segregated. The layer of ethyl acetate was placed in different flask. To clear up the ethyl acetate layer the method was repeated 3 times with ethyl acetate (3x400mL). In vacuum rotary evaporator, adjusting 100 rpm and 35°C temperature, ethyl acetate was evaporated. The extracted crude obtained have ethyl acetate which was weighed and kept in freezer at -20°C in a labeled flask.

Using 400mL of n-butanol, the water layer was further analyzed in the separating funnel. These two solvents were mixed and dissolved for 15 to 20 minutes. Pressure has to be released occasionally during mixing. Almost, 30 minutes were given to separating funnel in order to separate the layers. Collected n-butanol fraction in different flask and repeated the method twice more (2x400mL). To condense the n-butanol, it was evaporated at 100 rpm and 50°C temperature so that no further compound gets from water in n-butanol. After this, the extracted crude of n-butanol was weighed and kept in freezer at -20°C in a labeled flask.

Lastly, the water was evaporated at 60°C at about 100 rpm by using rotary evaporator to get aqueous fraction. Later, the crude residue of water was weighed, gathered in a tagged bottle and kept in freezer at -20°C.

**Antibacterial activity**

To find the biological activities, the methanolic extracts of the particular plants were exposed to antibacterial activity. The mechanism used to investigate antibacterial activity against selected microorganisms was agar well method.

**Bacterial strains**

Determining the antibacterial activity of both plant extracts, four bacterial strains were used in which three were gram negative like *Pseudomonas aeruginosa* (*P. aeruginosa*), *Escherichia coli* (*E. coli*), *Klebsiella pneumoniae* (*K. pneumoniae*) and one was gram positive namely *Staphylococcus aureus* (*S. aureus*).

**Preparation of bacterial culture**

Cultures of certain bacteria were formulated in nutrient broth medium at 120 rpm for twenty-four hours at 37°C incubated in a shaker. These cultures were kept with temperature of 5°C in refrigerator after the bacterial strains were developed [25].

**Evaluation of antibacterial activity**

**Agar well diffusion method**

**Principle**

Plant extracts containing antimicrobials are spread out into the medium and merge in a plate freshly seeded with test organisms. The resulting inhibition zones will be evenly circular as there will be a confluent lawn of growth. The diameter of inhibition zones was measured in millimeters (mm).
Reagents
Mueller Hinton agar medium (1L)
In this, 33g of Mueller Hinton agar medium was dissolved in 1000ml of distilled water. The dissolved medium was autoclaved at 121°C for 15 minutes. The autoclaved medium was well mixed and poured into petri plates while still molten.

Procedure
The agar well diffusion assay was utilized to assess the antibacterial activity of both plant extracts with few alterations [26]. To prepare agar well medium, the melted Mueller Hinton agar was poured in petri dishes and allowed to consolidate. After consolidation, 6mm width holes were made into agar and 10 𝜇L of the four fractions (water, ethyl acetate, chloroform and n-hexane extracts) of plant extracts were added.

To prepare 100mg/ml base of crude plant extracts for both plants, 100mg of plant extracts were dissolved in 1ml of DMSO. Furthermore, stock extract of 0.01ml of certain herbs was poured into each hole on the seeded medium and kept standing for an hour for appropriate diffusion. The dishes were incubated for twenty-four hours at 37°C and the resulting inhibition zones were determined.

Antifungal activity
To find the biological activities, the crude materials of particular plants were exposed to antifungal activity. The mechanism used to investigate antifungal activity against selected microorganism was agar well method.

Fungal strain
In the identification of antifungal activity of certain plant extracts, PBF-1 was used.

Preparation of fungal culture
Fungus was developed in Sabouraud broth for 72 hours at 28°C. The growth of fungus can be observed by cloudiness in the broth medium. For further analysis, the fungus culture was stored in the refrigerator at 3-6°C.

Evaluation of antifungal activity
Agar well diffusion method
For the determination of fungicidal influence of plant extracts, the inhibition of mycelial growth of the fungus is used and is noticed as strong inhibition (there is no progress of the fungus), incomplete inhibition (where fungus has less growth than the normal) and no inhibition (full growth of the fungus).

Reagent (Potato Dextrose Agar Medium)
To prepare potato infusion, 200g peeled and sliced potatoes were boiled in 1liter distilled water for 30 minutes. Decanting through filter paper. Add 10 grams dextrose and 10 grams agar powder and pasteurized the medium by autoclaving at 121°C for 15 minutes.

Procedure
Nearly 20ml PDA solution and 2ml of fractions (water, ethyl acetate, chloroform and n-hexane) of plant extracts were combined well and the mixture was poured into pre-sterilized petri plates under Laminar air flow. The centers of petri plates were used for fungal strain. Later, the plates were incubated at 25°C for 48 hours. After incubation, the results were documented as strong inhibition (if test fungus shows no progress), incomplete inhibition (if test fungus has less growth than the normal) and no inhibition (if full growth of the test fungus). For reference, control plate without plant extract was also preserved.

Results and discussion
According to latest study, the antifungal and antibacterial activities of C. leiocolacinus and H. griffithii were recorded against different strains of bacteria including S. aureus, K. pneumonia, P. aeruginosa, E. coli and fungal strain like PBF-1. These plant extracts indicated variable activities.

Antibacterial activity
The in vitro antibacterial activity of different fractions (water, chloroform, ethyl acetate and n-hexane) of C. leiocolacinus and H. griffithii was assessed by Agar well assay against four pathogenic bacteria. The bacteria involved both gram negative and
gram positive. On screening basis, it showed that all the four fractions of *H. griffithii* and *C. leiocalycinus* showed evident antibacterial activities against all the test pathogenic bacteria indicating different zones of inhibition (Table 1 & 2). The fractions of *C. leiocalycinus* was significantly active against all bacterial strains. Aqueous phase played highest activity against *K. pneumoniae* with zone of inhibition 10mm, moderate activity against *P. aeruginosa* with zone of inhibition 9mm and least activity against *E. coli* and *S. aureus* with zones of inhibition 6mm and 7mm respectively. Consequently, chloroform fraction demonstrated highest activity against *E. coli* with zone of inhibition 13mm, moderate activity against *K. pneumoniae* with zone of inhibition 9mm and least activity against *P. aeruginosa* and *S. aureus* with zones of inhibition 8mm and 7mm respectively. In addition, n-hexane phase demonstrated strong activity against *S. aureus* with zone of inhibition 7mm, moderate activity against *P. aeruginosa* and *E. coli* with zones of inhibition 5mm and 6mm and smallest activity against *K. pneumoniae* with the inhibition zone 3mm. Ethyl acetate phase exhibited strong activity against *K. pneumoniae* and *P. aeruginosa* with zones of inhibition 8mm for both, moderate activity against *S. aureus* with zone of inhibition 6mm and smallest activity against *E. coli* with zone of inhibition 5mm.

### Table 1. Zones of inhibition in millimeter of *C. leiocalycinus* against different bacterial strains

<table>
<thead>
<tr>
<th>Fractions</th>
<th>S. aureus</th>
<th>K. pneumoniae</th>
<th>E. coli</th>
<th>P. aeruginosa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>7</td>
<td>10</td>
<td>6</td>
<td>9</td>
</tr>
<tr>
<td>Chloroform</td>
<td>7</td>
<td>9</td>
<td>13</td>
<td>8</td>
</tr>
<tr>
<td>n-hexane</td>
<td>7</td>
<td>3</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>Ethyl acetate</td>
<td>6</td>
<td>8</td>
<td>5</td>
<td>8</td>
</tr>
</tbody>
</table>

The fractions of *H. griffithii* extracts exhibited activity to different extent against different bacterial strains. Aqueous fraction exhibited highest activity against *P. aeruginosa* and *E. coli* with zones of inhibition 10mm and 11mm, moderate activity against *S. aureus* with zone of inhibition 8mm and smallest activity was showed against *K. pneumoniae* with small inhibition zone 6mm. Furthermore, chloroform portion showed good activity against *K. pneumoniae* with zone of inhibition 10mm, moderate activity against *S. aureus* and *E. coli* with zones of inhibition 8mm and least activity was showed against *P. aeruginosa* with small zone of inhibition 6mm. Additionally, n-hexane part demonstrated good activity against *S. aureus* with zone of inhibition 9mm, least activity for *E. coli* with small zone of inhibition 4mm and did not show any activity against *K. pneumoniae* and *P. aeruginosa*. Consequently, ethyl acetate section showed good activity against *E. coli* with zone of inhibition 9mm, least activity against *K. pneumoniae* and *P. aeruginosa* with zones of inhibition 7mm. Whereas, ethyl acetate lacks the property by showing any activity against *S. aureus* (figure 1).

### Table 2. Zones of inhibition in millimeter of *H. griffithii* against different bacterial strains

<table>
<thead>
<tr>
<th>Fractions</th>
<th>S. aureus</th>
<th>K. pneumoniae</th>
<th>E. coli</th>
<th>P. aeruginosa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>8</td>
<td>6</td>
<td>11</td>
<td>10</td>
</tr>
<tr>
<td>Chloroform</td>
<td>8</td>
<td>10</td>
<td>8</td>
<td>6</td>
</tr>
<tr>
<td>n-hexane</td>
<td>9</td>
<td>N-A</td>
<td>4</td>
<td>N-A</td>
</tr>
<tr>
<td>Ethyl acetate</td>
<td>N-A</td>
<td>7</td>
<td>9</td>
<td>7</td>
</tr>
</tbody>
</table>

No inhibition zone: N.A
Antifungal activity
The effect of water, chloroform, ethyl acetate and n-hexane fractions of *C. leiocalycinus* and *H. griffithii* were investigated for in vitro antifungal activity against selected fungus by Agar well assay. The test fungus employed was PBF-1.

The fractions of *C. leiocalycinus* exhibited mild antifungal activity. Aqueous fraction showed partial inhibition (test fungus has less growth) while, ethyl acetate showed strong inhibition (no progress of test fungus) (figure 2). However, chloroform and n-hexane demonstrated no inhibition (full growth of test fungus).

Considering the case of *H. griffithii*, all its fractions showed no inhibition (full growth of the test fungus) against PBF-1 except aqueous fraction which showed partial inhibition (test fungus has less growth than the normal).

In this study, the obtained results showed that the selected plants were utilized to investigate their antibacterial and antifungal activity. We noticed that the fractions of *C. leiocalcinus* exhibited considerable antibacterial and mild antifungal activities against all tested bacterial and fungal strains. Chloroform fraction of *C. leiocalcinus* showed good antibacterial activity while other fractions of *H. griffithii* and *C. leiocalcinus* showed mild antibacterial activity. Ethyl acetate fraction of *C. leiocalcinus* showed strong inhibition (no progress of test fungus).

Overall antifungal activity of *H. griffithii* and *C. leiocalcinus* was not significant (table 3). The achieved results indicate that the plant extracts inhibited the growth of different microorganisms. Therefore, it showed that the plant extracts contain substances which prevent the growth of different microorganisms. At different concentrations, various researches have proved that the plant extracts prevent the development of different microorganisms [27]. Antibacterial activity of the plant extracts is considered due to the occurrence of flavonoids, tannins and alkaloids [28]. Some researchers also noticed that the presence of these different secondary metabolites in the plant extracts are responsible for antimicrobial effect [27]. Plant extracts are traditionally utilized to heal the wounds and used as ear drop in the treatment of earache and ear boils. They are also utilized to control dysentery and diarrhea [29].
Table 3. Antifungal activity of *H. griffithii* and *C. leiocalycinus*

<table>
<thead>
<tr>
<th>Plant samples</th>
<th>Fractions</th>
<th>PBF-1</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>C. leiocalycinus</em></td>
<td>Water</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Chloroform</td>
<td>−</td>
</tr>
<tr>
<td></td>
<td>n-hexane</td>
<td>−</td>
</tr>
<tr>
<td></td>
<td>Ethyl acetate</td>
<td>++</td>
</tr>
<tr>
<td><em>H. griffithii</em></td>
<td>Water</td>
<td>−</td>
</tr>
<tr>
<td></td>
<td>Chloroform</td>
<td>−</td>
</tr>
<tr>
<td></td>
<td>n-hexane</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Ethyl acetate</td>
<td>−</td>
</tr>
</tbody>
</table>

No Inhibition (−), Mild Inhibition (+), Strong Inhibition (++)

Figure 2. Antifungal activity of ethyl acetate fraction of *C. leiocalycinus*

**Conclusion**

The main purpose of this study is to identify the existence of antibacterial and antifungal activities in the preferred native medicinal plants of Balochistan like *C. Leiocalycinus* and *H. griffithii* in the province of Balochistan. In this examination, the antibacterial and antifungal activities of both the specific plants (*C. leiocalycinus* & *H. griffithii*) were investigated. As a result, they exhibited substantial antibacterial and mild antifungal activities. This indicated the significance of *C. leiocalycinus* and *H. griffithii* in the treatment of different diseases. The result against different microorganisms showed the curative capacity of *C. leiocalycinus* and *H. griffithii*. It can be concluded that the extracts of *C. leiocalycinus* and *H. griffithii* can represent a fundamental part in the field of medicines. Apart from this, it can also be examined as herbal sources in pharmaceutics and can be used in food industry.

**Authors’ contributions**

Conceived and designed the experiments: A Mengal, Samiullah & A Baqi, Performed the experiments: A Mengal & A Baqi, Analyzed the data: Samiullah, N Khan & A Rehman, Contributed materials/ analysis/ tools: Samiullah & N Khan, Wrote the paper: A Mengal, A Baqi & Samiullah.

**References**


