Original article

A study on predominance of keratinophilic flora in soil of Jaipur, India

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1. Introduction

Keratin is an insoluble fibrous and recalcitrant protein that is arranged into soft and hard state depending on their amino acids and sulfur contents. Both of Keratin construct fundamental ingredients of mammals, birds, and fishes in form of wool, feathers, hairs, nails, horns, and hooves. Keratin is densely packed and stabilized with a mechanical strength by certain hydrogen bonds, disulfide bonds, and hydrophobic interactions. Moreover, the cross-linking of protein chains hinders keratin degradation by common proteases (Hassan et al., 2020a; Fraser and Parry, 2003) (Figs. 1–5).

Keratinophilic fungi are imperfect molds which serve as saprophytic and consume their nourishment via decomposing the keratin or other organic matters (Kwon-Chung and Bennett, 1992; Rippon, 1988). Geographically, these are found in soils of forest, public places, marketplace, poultry sheds, herbivore or carnivore muck parks, as well as sediments of rivers and oceans containing humus and organic material, sewage and bird’s nest, barber’s hair dumping area (Kumawat et al., 2020; Gupta et al., 2012). In recent years, keratinophilic microbiota is attaining remarkable attention throughout the world (Sharma et al., 2020; Bohacz and Korniłowicz-Kowalska, 2019; Gupta et al., 2012; Bentubo et al., 2006) (Tables 1 and 2).

Therefore, a precise distribution and appearance of keratinophilic fungi help to view advantageous and unhealthy impacts on nature. As a point of industrial and ecological interests, these could possess a role in degradation of keratinous waste as well as production of Keratinolytic enzymes. The primary outcomes are of great interest in their utilization in a natural and environmentally friendly way. These may appropriate in fabrication of bioplastics; animal feed and fertilizer productions; biogas production; detergent activity to tolerate various surfactants, detergents, and organic solvents; dehairing in leather and textile industry (Hassan et al., 2020b; Bohacz and Korniłowicz-Kowalska, 2019). Although keratinophilic fungal appearance on earth manifests an emphatic impression, besides also designates a reservoir for primary infection. It may be at least for some pathogenic fungi or also...
Fig. 1. Prevalence of Keratinophilic Fungal Genera on Various Baits.

Fig. 2. Neighbor joining tree of isolated Arthroderma multifidum (KU578107) with reference data.

Fig. 3. Maximum Likelihood tree of isolated Arthroderma multifidum (KU578107) with reference data.
Table 1
Prevalence of Keratinophilic Fungal Genera on Various Baits.

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Name of fungal Genera</th>
<th>Human Hair</th>
<th>Human Nails</th>
<th>Chicken Feathers</th>
<th>Animal Hair</th>
<th>Total Plates</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Alternaria sp.</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>0</td>
<td>3</td>
<td>1.85</td>
</tr>
<tr>
<td>2.</td>
<td>Arthroderma sp.</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>3</td>
<td>1.85</td>
</tr>
<tr>
<td>3.</td>
<td>Aspergillus sp.</td>
<td>5</td>
<td>4</td>
<td>11</td>
<td>9</td>
<td>29</td>
<td>17.9</td>
</tr>
<tr>
<td>4.</td>
<td>Chaetomium sp.</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>4</td>
<td>2.47</td>
</tr>
<tr>
<td>5.</td>
<td>Chrysosporum sp.</td>
<td>8</td>
<td>2</td>
<td>8</td>
<td>6</td>
<td>24</td>
<td>14.81</td>
</tr>
<tr>
<td>6.</td>
<td>Emericella sp.</td>
<td>1</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>8</td>
<td>4.94</td>
</tr>
<tr>
<td>7.</td>
<td>Fusarium sp.</td>
<td>9</td>
<td>1</td>
<td>2</td>
<td>6</td>
<td>18</td>
<td>11.11</td>
</tr>
<tr>
<td>8.</td>
<td>Histoplasma sp.</td>
<td>4</td>
<td>1</td>
<td>4</td>
<td>6</td>
<td>15</td>
<td>9.26</td>
</tr>
<tr>
<td>9.</td>
<td>Malbranchea sp.</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>4</td>
<td>6</td>
<td>3.7</td>
</tr>
<tr>
<td>10.</td>
<td>Microsporum sp.</td>
<td>4</td>
<td>6</td>
<td>2</td>
<td>4</td>
<td>16</td>
<td>9.88</td>
</tr>
<tr>
<td>11.</td>
<td>Penicillium sp.</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>1.23</td>
</tr>
<tr>
<td>12.</td>
<td>Scopulariopsis sp.</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>5</td>
<td>3.09</td>
</tr>
<tr>
<td>13.</td>
<td>Trichophyton sp.</td>
<td>14</td>
<td>8</td>
<td>5</td>
<td>2</td>
<td>29</td>
<td>17.9</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>48</td>
<td>29</td>
<td>42</td>
<td>43</td>
<td>162</td>
<td></td>
</tr>
</tbody>
</table>
for those which are potentially pathogenic for small to big wild fauna (Garg et al., 1985). All keratinophiles endure as self-sufficient saprophytes in soil until the availability of favorable environmental conditions. Opportunistically or accidentally, being a contagion, these organisms adhere and penetrate to both humans and animal tissue through specific metabolic reactions and lead up to superficial and cutaneous fungal infections. Such keratinophiles are referred to as dermatophytes (Sharma et al., 2020; Bentubo et al., 2006). In the present study, the main objective was to evaluate prevalence of keratinophilic flora in soil of Jaipur (Rajasthan), India with their sustainable management.

2. Materials & methodologies

2.1. Soil collection

The selection of collecting soil sample areas for this study was based on the contamination of higher keratin levels. In the present study, a total of 50 soil samples was collected from various sites in Jaipur, India. After collection of soil samples, initial pH of soil was determined following the protocol of Rousk et al., 2009 using calibrated pH meter (Model 181, Electronics India).

2.2. Isolation and identification of keratinophilic fungi

Isolation of keratinophilic fungi was done according to Vanbreuseghem’s hair bait technique (Vanbreuseghem, 1952). After the visible appearance of mycelium on applied baits, a fungal colony was transferred to Sabouraud’s Dextrose Agar (SDA) medium (Hi-Media). After the purity of isolated fungal growth, fungus was identified on the basis of macroscopic characteristics. Subsequently, microscopic characteristics were also observed by flag scotch tape method at 100X power under microscope.

The cultural and morphological characteristics of fungal colonies and their identification were done by referring laboratory methods in basic mycology (Forbes et al., 2002), Descriptions of Medical Fungi (Kidd et al., 2016) and Pictorial Atlas of Soil and Seed Fungi (Watanabe, 1937). Keratinolytic activity of fungal isolates was determined following the protocol of Kacinova et al., 1989. On the basis of occurrence and highest keratinolytic activity, most prevalent fungi were selected for further studies.

2.3. Molecular identification

Whole-cell DNA from mycelial growth of isolate was extracted by following the protocol of Lee and Taylor (1990). The quality of extracted DNA was monitored by agarose gel electrophoresis using 1.5%agarose gel with ethidium bromide fluorescence. ITS (Internal transcribed spacer) regions of fungal DNA were amplified using ITS primer such as ITS1(5'-CTCTGAGTGGAACCTTGCGG-3') and ITS4 (5'-TCCGCTATTGATATGC-3') respectively (Sigma-Aldrich, Bengaluru, India) by Eppendorf's DNA gradient thermal cycler (White et al., 1990; Sharma et al., 2017). Sequencing reactions were carried out using Big Dye terminator cycle sequencing kit, version3.1 (Applied Biosystems, CA, USA) and analyzed by the ABI3130 genetic analyzer (Applied Biosystems, CA, USA).

The resultant sequence was compared with the GenBank data base using NCBI Basic Local Alignment Search Tool (BLAST). The sequence showing > 99% match with data base sequences of the reference taking as for species identification. The sequences were aligned with Clustal-ω computer program in MEGA-10 software. Phylogenetic trees were prepared by the Neighbor-Joining and Maximum Likelihood methods using MEGA10 (Kumar et al., 2018).

2.4. Anti-fungal assay

The anti-fungal activity against selected test fungi was evaluated by disc diffusion (Gould and Bowie, 1952) method. Essential oils were extracted from Ocimum tenuiflorum (Tulsa), Citrus grandis (Pomelo), and Eucalyptus globulus (Bluegum) plants. In assay, sterilized discs impregnated with 10 μL of crude essential oil extracted from respective plants were aseptically transferred onto SDA plates inoculated with test fungi. The plates were then incubated at 30 °C for 48–72 h and were observed for the formation of zone of inhibition. The activity index was also calculated by following formula. Activity index(AI) = Inhibition Zone by Essential Oil / Inhibition Zone of Standard

Table 2

Antifungal Activity of Essential Oils Extracted from Plants

<table>
<thead>
<tr>
<th>Test Fungi</th>
<th>Ocimum tenuiflorum</th>
<th>Eucalyptus globulus</th>
<th>Citrus grandis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Inhibition Zone (IZ)</td>
<td>Activity Index (AI)</td>
<td>Inhibition Zone (IZ)</td>
</tr>
<tr>
<td>Arthrodema multifidum</td>
<td>20.5 ± 1.75</td>
<td>0.72</td>
<td>23.5 ± 1.32</td>
</tr>
<tr>
<td>Chrysosporium indicum</td>
<td>20.75 ± 1.66</td>
<td>1.46</td>
<td>17.5 ± 0.5</td>
</tr>
<tr>
<td>Fusarium oxysporum</td>
<td>16.75 ± 0.52</td>
<td>1.08</td>
<td>16.5 ± 0.50</td>
</tr>
</tbody>
</table>

Note: IZ = Inhibition Zone (in mm) with the diameter of disc (6 mm), AI = Activity Index.

2.5. Statistical analysis

All Experiments were performed in triplicate and data analyzed are mean ± SE subjected to one-way ANOVA with significant (P < 0.05).

3. Results and discussion

In pH analysis of soil samples, maximum pH 10.6 was recorded for soil samples collected from Sambhar Lake site following by soil samples collected from dump sites of barbers shops which were exhibited more alkaline pH between pH 7.9–9.55. Likewise, other sites: road side, poultry farms, cattle yard, and public places of Jaipur city were found to have a pH range of 6.5–8.95. In the present study, numerous mycelium structures were observed as hyphae and spores which were recorded to be...
keratinophilic fungi. Total twenty-four species distributed in thirteen genera were isolated as keratinophilic and related fungi. *Chrysosporium* sp. (14.81%) was second most dominant keratinophilic fungi afterward to *Trichophyton* sp. (17.90%) and *Aspergillus* sp. (17.90%) and the least account was recorded for *Penicillium* sp. (1.23%) (Figure-1 & Table-1). The major incidence ratio of keratin degradation in baiting technique was obtained on Human Hair > Animal Hair > Chicken Feathers > Human Nails (high to low growth) baits.

According to results, growth of *Malbranchea saccardo* (3.70%), *Chaetomium kunze* (2.47%) and *Arthroderma multifidum* (1.85%) were reported for the first time in Jaipur, Rajasthan. The results of present study are in line with several researchers who have documented the distribution of keratinophilic fungi from soil of India (*Randhawa and Sandhu, 1965; Deshmukh, 1999; Vidal, 2000*). Pakshir et al. (2013) also reported that most (66.42%) of the keratinophilic fungi grow in the soil with pH 7.0–8.0. Findings of Kumawat et al., 2020 are in support of the present study that has reported 154 isolates belonging to 16 genera and 31 species and recovered to *Chrysosporium tropicum* as most predominant fungal species in the soil of Rajasthan.

In molecular identification, DNA was isolated using the phenol-chloroform method and its purity was confirmed to be between 1.9 and 2.6 at 260 nm/280 nm ratios by using NanoDrop™ 2000/2000c Spectrophotometers. The PCR amplification of ITS region of the isolate KU7 and KUB yielded PCR products of about 550 bp, respectively. Ramaraj et al. (2016) obtained the amplicon size of 850–800 bp for amplified regions (ITS-1 and ITS-2) of keratinophytes using primers ITS-1 and ITS-4 that is similar to our findings.

The PCR product with primer pair ITS1 was sequenced for species identification and found to be 90 to 100% similar in the BLAST program to sequences of the ITS1, 5.8S rRNA gene, and ITSII regions of respective fungi. On the basis of BLAST program results, ITS1 sequence of isolates was determined as *Arthoderma multifidum* (KU578107) and *Chrysosporium indicum* (KU578108) which was similar to morphological identification. After a preliminary identification, sequences of *A. multifidum* and *C. indicum* were deposited in the GenBank database (http://www.ncbi.nlm.nih.gov/Genbank/index.html) with the accession numbers KU578107 & KU578108 respectively.

The multiple sequence alignment analysis of isolated *A. multifidum* was found to similar with reference AB861744.1 and AB861799.1 at GenBank data. There were a total of 547 positions for *A. multifidum* in final dataset.

The data of thirteen fungal sequences including one test sequence and twelve reference sequences were used in phylogeny construction. In the phylogenetic tree, *A. multifidum* found to associate with *A. multifidum*, *C. keratinophilum*, *C. articularatum*, *A. terreus*, *Aphanoascus fulvescens*, *A. verrucosus*, *A. reticulisporus*, *C. tropicum*, *C. zonatum*, *A. clathratus*, *A. arxii*, and *Uncinocarpus orissiari* members in ITS1 homology group (Figure-1).

In the Neighbor-Joining tree of isolated *Arthoderma* sp. strain, the branch length obtained with the sum of 16.95 branch lengths. In NJ tree, there are two major clusters A & B. The members in cluster A were classified into two groups with ITS1 homology (groups a and b) according to their ITS1 DNA sequences. In Cluster A, test KU578108.1 and reference (AJ439446.1) sequence of *C. indicum* were found to have very close relationships with *Aphanoascus* sp. (AJ439440.1&AJ439434.1). Subsequently, in Cluster B, and group b of cluster A, maximum members were recorded for *Chrysosporium* sp. with 100% bootstrap support. *C. indicum* was shown to be closely allied with the anamorphs species *A. durus* and is on a well-supported branch that contains another major lineage with another as *A. punsolar* lineage. The phylogenetic relationships mentioned above were also supported by the ML tree where the sequence of *C. indicum* (KU578108.1) was found to be closely allied with reference sequence AJ439446.1 with the highest log-likelihood of −9547.26. This species was also grouped together with *Trichophyton* anamorphs of *Arthroderma* sp. by Vidal, 2000.

Results of present study have indicated about phylogenetic relationships between the teleomorphic and anamorphic form of keratinophilic fungi and exhibited a reason for phase changes between geophilic and dermatophytic fungi by accident or opportunistic where they may cause infections in both humans and animals on the opportunity and may exist as parasite on the body. In this dimorphism character of fungi, a number of fungal (superficial and deep mycotic) infections has enhanced as a significant clinical problem and still is continued (NCCLS, 1997; Ataide et al., 2011). The limited therapeutic strategies, ranging from routes administration, efficacy, cost, solubility and stability of antifungal agents needed a search of novel antifungal agents from biosynthetic laboratories i.e. plants (Sharma et al., 2014).

In this approach, anti-fungal activity was used as research parameter to control incidence of fungal infections and recorded that essential oil extracted from all selected plants were found to have great antifungal potential against all keratinophilic fungal species tested. However, a comparison showed that essential oil of *Eucalyptus globulus* has a greater potential of anti-fungal activity than those of other essential oil. The *Eucalyptus globulus* exhibited highest zone of inhibition against *A. multifidum* with an activity index of (IZ: 23.5 mm; AI: 0.82). In support, essential oil of *Ocimum tenuiflorum* was found to exhibit highest zone of inhibition against *Chrysosporium indicum* with an activity index of (IZ: 20.75 mm; AI: 1.46). *Citrus grandis* was also found to exhibit its highest zone of inhibition against *A. multifidum* with an activity index of (IZ: 12.62 mm; AI: 0.44). In respect of Fusarium oxysporum, fungal
growth was found to inhibit by essential oil of *Ocimum tenuiflorum* with an activity index of (IZ: 16.75 mm; AI: 1.08).

In Minimum inhibitory concentration (MIC), range 1/7(1/4) for all oil *Ocimum tenuiflorum* and *Eucalyptus globulus* was found to be effective for all tested fungi excluding to *Citrus grandis*, which was found to be effective in more decreased concentration.

The present results are in line with Basilio and Basilio, 1999 who reported the successful antifungal activity of *O. basilicum* oil against *Aspergillus ochraceus* (Basilico and Basilio, 1999). Vasudeva and Sharma (2012) reported maximum antifungal activity for essential oil extracted from *Citrus species* against *Fusarium oxysporum* with zone of inhibition 10.23 mm.

4. Conclusion

In the present study, the more prominent occurrence of Keratinophilic fungi in soils of Jaipur District confirms semi-arid environment of city. Certain reservoirs can part in approaching to superficial infections such as tinea or ringworm via frequent human contact under favorable conditions. The leading incidence of keratinophilic fungi generates an urgent need to manage the appearances of fungi. Results obtained regarding antifungal activity prove to be useful and may replace expensive therapies for controlling fungal infections and provide affordable remedial resources for millions of individuals or poor people around the world.

Although, the isolated keratinolytic fungi are also facilitated the breakdown of protein and di-sulphide bonds in keratin which could be exploited for many industrial applications. The crude keratinase enzyme could also be employed as an effective and eco-friendly alternative in leather processing industries. The results of this research work can guide and lead to the anticipated path for biotechnological interventions.

Acknowledgement

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