Morpho-Molecular Identification of
*Trichoderma asperellum* CA-03/9840

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Most isolates of the genus *Trichoderma* were found to act as mycoparasites of many economically important aerial and soil-borne plant pathogens. *Trichoderma* has gained importance as a substitute for chemical pesticides and hence an attempt was intended to corroborate the positive relatedness of molecular and morphological characters. A fungal strain of *Trichoderma asperellum CA-03/9840* was isolated from a soil sample collected from Farmer Field, Sultanpur, Uttar Pradesh, India. The universal primers were used for amplification of 5.8SrRNA gene fragment and the strain was characterized by using 5.8SrRNA gene sequence with the help of ITS marker. It is proposed that the identified strain *Trichoderma asperellum CA-03/9840* be assigned as the type strain of a species of the genus *Trichoderma* based on TrichoKey analysis together with the 5.8SrRNA gene sequence search in Ribosomal Database Project, small subunit rRNA and large subunit rRNA databases. The sequence was deposited in GenBank with the accession number KU821782. Thus an integrated approach of morphological and molecular markers can be employed to identify a superior strain of *Trichoderma* for its commercial exploitation.

**Keywords:** 5.8S ribosomal RNA gene, *Trichoderma*, ITS, Biocontrol, Antagonist.

*Trichoderma*, commonly available in soil and root ecosystem, has gained immense importance since last few decades due to its biocontrol ability against several plant pathogens. Some strains of *Trichoderma* like *T. harzianum*, *T. atroviride*, *T. viride*, *T. virens* and *T. koningii* are efficient bio control agents which have the ability to inhibit pathogen growth in the soil, hence improving the overall health of the plant. Antagonistic microorganisms such as *Trichoderma* reduce growth, survival of pathogen by different mechanism like competition antibiosis, mycoparasitism, hyphal interactions and enzyme secretion. Such micro organisms are now available commercially and are used in crop management and practices Kubicek, et. al. (2000). The use of *Trichoderma* species as biological control agents has been investigated for over 70 years but it is only relatively recently that strains have become available commercially. Biocontrol agents are widely regarded by public as “natural” and non-threatening products, although risk assessments must clearly be carried out on their effects on non-target organisms. Moreover, knowledge concerning the behaviour of such antagonists is essential for their effective use. The morphological and microscopic figure of *Trichoderma asperellum* CA-03/9840 is given figure 1 (a-c).

*T. asperellum* was described as a new species in 1999 for isolates that produce finely warted conidia, with temperature optima of 30°C.
and survive by producing chlamydospore. It is used as biological control agent against a wide spectrum of plant disease-causing organisms, including straminopiles such as Phytophthora megakarya fungi and nematodes. The species also has been shown to have antibacterial activity through the production of trichotoxin peptaibols.

**Habit and Habitat**

Isolate Name: T. asperellum CA-03

Isolated from: Sultanpur Field Chilli

ITCC No.: 9840

Accurate and definitive fungal identification is essential for correct disease diagnosis, treatment of associated with fungal infections. Characterization of fungal species using classical methods is not as specific as the genotyping methods. Genotypic techniques involve the amplification of a phylogenetically informative target, such as the small-subunit (5.8S) rRNA gene reported by Woese CR et al. (1977). rRNA is essential for the survival of all cells, and the genes encoding the rRNA are highly conserved in the fungal and other kingdoms. The sequences of the rRNA and proteins comprising the ribosome are highly conserved throughout evolution, because they require complex inter- and intramolecular interactions to maintain the protein-synthesizing machinery reported by Sacchi CT, et. al. (2002), Hillis DM, et. al. (1991) and Woese CR. et. al (1987). *Trichoderma* species are common soil inhabitants and are effective in providing biocontrol of soil borne pathogens due to antagonistic behaviors. The major aspect of successful biological control strategies includes the production, formulation and delivery system of bio-agents. The internal transcribed spacer (ITS) region of the rDNA is perhaps the most widely sequenced DNA region in fungi. It has typically been most useful for molecular systematic study at species level, and even within species found by Ospina-Giraldo MD, et. al. (1998), Kubicek CP et al. (2000), Kulling-Gradinger CM et. al. (2002) and Lee C et. al (2002) attempted a first phylogenetic analysis of the whole genus of *Trichoderma* using sequence analysis of the ITS region of rDNA.

In this study, the method of isolation and identification of an unknown fungal from Farmer Field of Sultanpur district using 5.8SrRNA gene

**Morphological Key**

<table>
<thead>
<tr>
<th>Morphological Observation</th>
<th>Species Character</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colony growth rate (cm)</td>
<td>7-8 cm</td>
</tr>
<tr>
<td>Colony colour</td>
<td>Dark yellowish green</td>
</tr>
<tr>
<td>Reverse colony colour</td>
<td>Colourless</td>
</tr>
<tr>
<td>Colony edge</td>
<td>Smooth</td>
</tr>
<tr>
<td>Culture smell</td>
<td>No characteristic odour</td>
</tr>
<tr>
<td>Mycelial form</td>
<td>Smooth</td>
</tr>
<tr>
<td>Mycelial colour</td>
<td>Cottony white</td>
</tr>
</tbody>
</table>

**Cultural characteristic**

<table>
<thead>
<tr>
<th>Cultural Character</th>
<th>Species Character</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conidiation</td>
<td>Ring like zones</td>
</tr>
<tr>
<td>Conidiophore branching</td>
<td>Branched, regular</td>
</tr>
<tr>
<td>Phialide shape</td>
<td>Cylindrical</td>
</tr>
<tr>
<td>Phialide size</td>
<td>6.2-10.5 x 3.1-3.9 m</td>
</tr>
<tr>
<td>Conidial shape</td>
<td>Globose to subglobe</td>
</tr>
<tr>
<td>Philospore size</td>
<td>2.6-3.0 x 2.0-2.4 m</td>
</tr>
<tr>
<td>Conidial wall</td>
<td>Smooth</td>
</tr>
<tr>
<td>Conidial colour</td>
<td>Light green</td>
</tr>
<tr>
<td>Chlamydospore</td>
<td>present</td>
</tr>
<tr>
<td>Spore Germination time</td>
<td>12 -13 Hours</td>
</tr>
</tbody>
</table>

Fig. 1. (A-C): *Trichoderma asperellum CA-03/9840* strain in PDA medium and microscopic image.
sequence as reported in bacterial rRNA gene found by Srivastava S, et. al. (2008) to characterize the strain CA-03/9840 as a member of the *Trichoderma* spp. The soil sample has received great attention from the public, due to its potential for biodiversity and biological conservation. The internal transcribed spacer (ITS) region of the rDNA is perhaps the most widely sequenced DNA region in fungi. It has typically been most useful for molecular systematic study at species level, and even within species Kindermann et al. (1998) attempted a first phylogenetic analysis of the whole genus of *Trichoderma* using sequence analysis of the ITS1 region of rDNA.

**MATERIALS AND METHODS**

**Isolation and identification of *Trichoderma***

Soil samples were collected from various experimental fields of Indian Farmer Field of Sultanpur district. Isolate of *Trichoderma* species was isolated and identified in potato dextrose agar (PDA) with low sugar medium reported by Hiney M, et. al. (1992). The identification of *Trichoderma* isolates were confirmed both by morphological and molecular characters (ITS) NCBI, GenBank Accession Number KU821782 and reconfirmed by Indian Type Culture Collection (ITCC), IARI, Pusa, New Delhi Accession Number allotted 9840.

**DNA isolation of *Trichoderma***

Pure culture of the target fungal was grown overnight in liquid PD Broth medium for the isolation of genomic DNA using a method described by Hiney et. al. (1998).

**Molecular characterization**

The total genomic DNA was extracted from isolate of *Trichodermaasperellum*CA-03/9840based on CetrimeTetradecylTrimethylAmmonium Bromide (CTAB) mini extraction method of Crowhurst et al. (1995) with minor modification.

**Agarose gel electrophoresis**

Ten microlitre of the reaction mixture was then analyzed by submarine gel electrophoresis using 1.0% agarose with ethidium bromide at 8 V/cm and the reaction product was visualized under Gel Doc/UV trans-illuminator.

**Internal Transcribed Spacer region**

The internal transcribed spacer (ITS) regions of the rDNA repeat from the 3’end of the 5.8S and the 5’end of the 5.8S gene were amplified using the two primers, ITS-1 and ITS-4 which were synthesized on the basis of conserved regions of the eukaryotic rRNA gene reported by Zhang Z, (2000). The PCR-amplification reactions were performed in a 50 ml mixture containing 50 mM KCl, 20 mM Tris HCl (pH 8.4), 2.0 mM MgCl2, 200 mM of each of the four deoxynucleotide triphosphates (dNTPs), 0.2µM of each primer, 40 ng/ml of template and 2.5 U of Taq polymerase. The cycle parameters included an initial denaturation at 94°C for 5 min, followed by 40 cycles of denaturation at 94°C for 1 min, primer annealing at 55°C for 2 min and primer extension at 72°C for 3 min and a final extension for 10 min at 72°C. Amplified products were separated on 1.2% agarose gel in TAE buffer, pre-stained with ethidium bromide (1mg/ml) and electrophoresis was carried out at 60 volts for 3 hours in TAE buffer. One Kb ladder (MBI, Fermentas) was used as a marker. The gel was observed in a trans-illuminator over ultra violet light. The desired bands were cut from the gel with minimum quantity of gel portion using QIAGEN gel extraction kit.

**Purification of PCR product**

The PCR product was purified by Qiagen gel extraction kit using the following protocol described below. The DNA fragment was excised from the agarose gel with a clean sharp scalpel. Then the gel slice was weighed in an eppendorf. We then added 3 volumes of buffer QG to 1 volume of gel (100 mg ~ 100 ml). The mixture was then incubated at 50°C for 10 min. The gel was dissolved by vortexing the tube every 2 to 3 min during the incubation until the mixture color is uniformly yellow. We then added 1 gel volume of isopropanol to the sample and mixed. A QIAquick spin column is then placed in a 2 ml collection tube provided. The sample is applied to the QIAquick column followed by centrifugation for one minute so that DNA binds to the column. The flow-through is discarded and the QIAquick column is placed back in the collection tube. We then added 0.75 ml of buffer PE to QIAquick column and centrifuged for 1 min to wash. The flow through is again discarded and the QIAquick column centrifuged for an additional 1 min at 10,000 x g. The QIAquick column is now placed into a clean 1.5 ml eppendorf. We then added 50ml of buffer EB (10 mM Tris-Cl, pH 8.5) to the center of the QIAquick membrane and...
Fig. 2. TrichoKey analysis of *Trichoderma asperellum* strain CA-03
centrifuged the column for 1 min to elute DNA.

**DNA sequencing of the 5.8SrDNA fragment**

The 5.8SrDNA amplified PCR product (100 ng concentration) was used for the sequencing with the single 5.8SrDNA 20F Forward, ITS1 primer: 5'- TCCGTAGGTGAACCTGCGG -3' and 22R Reverse ITS4 primer: 5'- TCCTCCGCTTAT TGATATGC -3' synthesized by DNA Sequencer by (Merck laboratory, Bangalore).

**Sequence Analysis**

A comparison of the 5.8SrRNA gene sequence of the test strain was done using BLAST against non-redundant nucleotide (nr/nt) database observed by Thompson JD et al. (1994). A number of *Trichoderma* sequences were selected on the basis of a similarity score of 90% with database sequences. Multiple sequence alignment of these selected homologous sequences and 5.8SrRNA gene sequence of test strain was performed using Clustal W reported by Saitou N, et al. (1987). Subsequently, an evolutionary distance matrix was then generated from these nucleotide sequences in the dataset. A phylogenetic tree was then drawn using the Neighbor Joining method reported by Tamura K, et al. (2007). Phylogenetic and molecular evolutionary analyses were conducted using MEGA (Molecular Evolutionary Genetics analysis) version 4.0 reported by Altschul SF, et al. (1997). We again compared the 5.8SrRNA gene sequence of test strain with different set of sequence database such as small subunit ribosomal RNA (SSUrRNA) and large subunit ribosomal RNA (LSUrRNA) using Ribosomal RNA BLAST reported by Cole JR, (2009). 5.8SrRNA gene sequence of test strain is also compared against those sequences, in Ribosomal Database Project found by Wang Q, et al. (2007) by using the RDP Classifier check Program. The annotated information for the sequence in the database to which 5.8SrRNA aligns is used for the fungal identification.

**RESULTS**

A total of 573bp of the 5.8SrRNA gene was sequenced and used for the identification of isolated fungal strain. Subsequently, 5.8SrRNA gene sequence based phylogenetic tree showing relationships between the test strain CA-03/9840 and selected representatives of the genus *Trichoderma* is given in Figure 1. It is evident from phylogenetic analysis of 5.8SrRNA gene that the isolate CA-03/9840 represents a genomic species in the genus *Trichoderma*. Comparison of test strain against known sequences of SSUrRNA and LSUrRNA databases showed that the gene sequence of isolate CA-03/9840 has 90% sequence similarity (Score=546 bits, Expect=0.0) with 5.8SrRNA gene sequence of *Trichoderma* (Genbank Acc. No.: KU821782). Thus, data shows that the isolate CA-03/9840 is a member of the genus *Trichoderma*. Similarity rank program classifier available at the Ribosomal Database Project classified the isolate CA-03/9840 as a novel genomic species of the genus *Trichoderma* with a confidence threshold of 90% (Figure 2). The 5.8SrRNA gene sequence of isolate CA-03/9840 was deposited in GenBank and allotted the accession number KU821782.

**DISCUSSION**

**Molecular analysis using Internal Transcribed Spacer region (ITS)**

Ribosomal RNA (rRNA) sequence analysis has been well-documented as a means of determining phylogenetic relationships in all of the major organismal domains. Variable sequences within the mature small subunit (SSU) and large-subunit (LSU) rRNA genes have been found to be appropriate for assessing subgeneric relationships in many eukaryotes. One of these variable regions, the D2 region of the LSU, has been used to determine phylogenetic relationships in a number of pathogenic fungal genera found by Logrieco, A, et al. (1995). The ITS region of the rDNA operon, which is more variable than the D2 region, has proven useful in distinguishing relationships at the species level found by Kusaba, M et. al. (1995).

The genetic variability within 69 biocontrol isolates of *Trichoderma* collected from different geographic locations and culture collections and their phylogenetic analysis were done with the help of the sequence data obtained from the inter Transcribed spacer 1 (ITS1) region of Ribosomal DNA and a fragment of the translation elongation factor 1 (tef1) and reported that more than 50% of the potential bio-control strains were...
grouped within *Trichoderma* section Pachybasium reported by Hermosa, M. et al. (2004).

*Trichoderma* isolates with different biocontrol capabilities and identification by molecular methods and further characterized into three main clades by internal transcribed spacer (ITS) Sequence analyses. Consequently, a reliable phylogenetic tree was constructed containing isolates belonging to the *T. harzianum* clade reported by M. Maymon et al. (2004).

Molecular phylogenetic analyses of biological control strains of *Trichoderma* (*Ascomycetes, Hypocreales*) strains that have warted conidia are traditionally identified as *T. viride*, the type species of *Trichoderma*. However, two morphologically distinct types of conidial warts (I and II) have been found. Because each type corresponds to a unique mitochondrial DNA pattern, it has been questioned whether *T. viride* comprises more than one species. Combined molecular data (sequences of the internal transcribed spacer 1 [ITS-1] and ITS-2 regions and of part of the 5.8SrRNA gene along with results of restriction fragment length polymorphism analysis of the endochitinase gene and PCR fingerprinting), morphology, physiology, and colony characteristics distinguish type I and type II as different species. Type I corresponds to "true" *T. viride*, the anamorphic of *Hypocrea rufa*. Type II represents a new species, *T. asperellum*, which is, in terms of molecular characteristics, close to the neotype of *T. hamatum*.

Analysis of ITS1-5.8S-ITS2 region of the cDNA showed that approximate 600 bp and size variation was observed. Restriction analysis of this region showed that inter and intra-specific polymorphism found by Latha J et al. (2002).

*Trichoderma* has attained importance for substitute of chemical pesticides and hence an attempt was intended to corroborate the positive relatedness of molecular and morphological characters. A fungal strain of *Trichoderma longibrachiatum* 28CP/7444 was isolated from a soil sample collected from Barabanki district of Uttar Pradesh, India. The universal primers were used for amplification of the 28S rRNA gene fragment and strain characterized by using 28SrRNA gene sequence with the help of ITS marker. It is proposed that the identified strain *Trichoderma longibrachiatum* 28CP be assigned as the type strain of a species of the genus *Trichoderma* based on phylogenetic tree analysis together with the 28S rRNA gene sequence search in Ribosomal Database Project, small subunit rRNA and large subunit rRNA databases. The sequence was deposited in GenBank with the accession number JX978541. Thus, an integrated approach of morphological and molecular markers can be employed to identify a superior strain of *Trichoderma* for its commercial exploitation. Previously similar results were also reported by Shahid M et al. (2013, 2014) and they also concluded that most of the *Trichoderma* species are morphologically very similar and were considered as a single species for many years. Since new species were discovered, a consolidated taxonomical scheme was needed, proposed and defined nine morphological species aggregates.

DNA methods brought additional valuable criteria to the taxonomy of *Trichoderma* which are being used today for studies that include identification and phylogenetic classification.

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