

Potential antagonism of some *Trichoderma* strains isolated from Moroccan soil against three phytopathogenic fungi of great economic importance

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(Reçu le 05/04/2017; Accepté le 15/04/2017)

Abstract

In this study, 17 *Trichoderma* spp. strains were isolated from different soils (Argan forests and crop fields) in Morocco. Purified monospore cultures were identified using molecular methods and tested for their potential antagonism against three phytopathogenic fungi (*Fusarium oxysporum*, *verticillium dahlia* and *rhizoctonia solani*). After DNA extraction, translation elongation factor (tefl) was amplified in extracts of 17 strains, sequenced and compared with their ex-types. As a result, three species were identified among the strains, which clustered in two different subclades of *Trichoderma*: the species *T. afroharzianum*, and *T. guizhouense* belong to *Harzianum* clade and *T. reesei* species belongs to *Longibrachiatum* clade. Dual culture plate assay was conducted confronting 17 *Trichoderma* isolates against *Fusarium oxysporum*, *Verticillium dahlia* and *Rhizoctonia solani*. *In vitro*, *Trichoderma* isolates showed effective antagonistic activity by decreasing mycelium growth of soil borne pathogens. *Trichoderma afroharzianum* showed the highest Percentage of Inhibition of Radial Growth (PRIG %). The highest PRIG% = 98% is when 8A2.3 isolate confront *R. solani* and the lowest PRIG% = 67% is when T9i10 confront *F. oxysporum*. On the other hand, T9i12, which is *reesei* species, significantly inhibit mycelium growth of the three pathogens.

Keywords: *Trichoderma* spp, *Fusarium oxysporum*, *Verticillium dahlia*, *Rhizoctonia solani*, antagonistic fungi.

Résumé

Dans cette étude, 17 souches de *Trichoderma* ont été isolées à partir d'échantillons du sol de différentes origines (champs et forêt d'arganier). Les espèces ont été identifiées par la caractérisation moléculaire des cultures monospores des isolats de *Trichoderma* et leur potentiel d'antagonisme a été évalué contre trois champignons phyto-pathogènes telluriques (*Fusarium oxysporum*, *verticillium dahlia* et *rhizoctonia solani*). Après l'extraction de l'ADN, le facteur d'élongation et de traduction tefl a été utilisé comme marqueur moléculaire pour amplifier, séquencer et par la suite caractériser les 17 souches qui ont été comparées avec leur ex-types déjà référencés. Par conséquent, l'identification moléculaire par tefl a permis la caractérisation de trois espèces de *Trichoderma* à savoir *T. afroharzianum*, et *T. guizhouense* appartenant à la branche *Harzianum*, et *T. reesei* appartenant à la branche *Longibrachiatum*. Les 17 isolats candidats ont été sélectionnés par une méthode de criblage basée sur la PCR (polymerase chain reaction). L'évaluation du potentiel antagoniste des souches de *Trichoderma* contre les champignons phytopathogènes telluriques (*F. oxysporum*, *R. solani* et *V. dahliae*) a été réalisée par la méthode de confrontation sur boîtes de pétrie. L'étude *in vitro* de l'activité antagoniste par le «Pourcentage of Radial Inhibition Growth» (PRIG %) a permis de suivre l'évolution de la croissance radiale du mycélium des différents champignons phytopathogènes. Le pourcentage le plus élevé (PRIG% = 98%) a été observé pour la confrontation entre l'isolat 8A2.3 et *R. solani* et le pourcentage le plus faible (PRIG% = 67%) a été observé pour la confrontation entre l'isolat T9i10 et *F. oxysporum*. D'un autre côté, T9i12 qui est un isolat de l'espèce *T. reeseia* a engendré une inhibition élevée de la croissance radiale du mycélium des différents pathogènes.

Mots clés: *Trichoderma* spp, *Fusarium oxysporum*, *Verticillium dahlia*, *Rhizoctonia solani*, champignons antagonistes.

INTRODUCTION

Soil borne pathogens attack a wide range of susceptible plants and cause telluric diseases like seed decay, damping off, root rot and blights (Sneh et al., 1991; El Amraoui et al., 2015).

Some soil borne fungi survive for several years in soil because of their conserved structures then germinate and develop under favorable environmental conditions. In fact, management of diseases caused by soil borne pathogens is difficult. Synthetic fungicides and fumigants, though still used, are not considered a potential solution to sustainable plants production because of their non specific targets application and negative impact on human health and environment (James et al., 1992; Howard et al., 1994).

Trichoderma spp. is a cosmopolitan fungus common in different biotopes and shows different interactions with

their neighboring lives. *Trichoderma* spp. is abundant in soil and involves different kind of interaction with other micro-organisms in the rhizosphere. Some *Trichoderma* spp. are opportunistic and show parasitic lifestyle against other soil borne fungi what makes them interesting myco-parasitic fungi for the biocontrol of soil borne pathogens. Therefore, many researches in the literature recognized tremendous antagonistic profiles of different *Trichoderma* spp. against soil borne pathogens. These *Trichoderma* are used as biological control agents against fungal phytopathogenes (Chet, 1987; Cook, 1993; Papavizas, 1985; Dighton et al., 2005).

Selection of antagonistic candidates in this genus depends on their relevant antimicrobial capacity to reduce either pathogen population or perturb its pathogenic pathway (Rodrigez et al., 2008).

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Some *Trichoderma* species including *Trichoderma harzianum*, *Trichoderma atroviride*, *T. virens*, and *Trichoderma asperellum* revealed mycoparasitic mode of action. That is, almost all *Trichoderma* species showed necrotrophic hyperparasitism up on other fungi. The former synthesize chitinases and other proteases that degrade pathogenic fungi cells fed by *Trichoderma* (Elad and Kapat, 1999; Pozo et al., 2004; Sanz et al., 2005).

Ability of *Trichoderma spp.* to interact with and antagonize other fungal soil borne pathogens tightly depends on its biodiversity and natural habitat. In fact, *Trichoderma* species and strains contain tremendous antagonistic profiles due to production of a wide range of secondary metabolites and enzymes. Therefore, each isolate perform specific interaction at the site of living (Hyde, 2005; Suryanarayanan and Hawksworth, 2005).

Trichoderma sp. is known as producer of destructive membrane enzyme (i.e. chitinases) and used as commercial biocontrol agent. However, not all of *Trichoderma* species are potential antagonists. Therefore, molecular identification of species belonging to *Trichoderma* genus is mandatory. In fact, molecular markers are widely used during screening of *Trichoderma* antagonistic candidates to avoid any misleading in identification, monitoring and/or labeling of potential antagonistic candidates (Watts et al., 1988; Schmoll, 2014; Jaklitsch, 2015).

ITS1 and ITS2, *tefl* and/or RNA polymerase gene *rpb2* are the most used molecular markers in phylogenetic analysis for identification and characterization of *Trichoderma* species (Jaklitsch, 2006; Jaklitsch, 2011).

In addition, this molecular tool has been developed for high throughput sensitive identification of *Trichoderma spp.* and considered as a crucial step in early screening of potential antagonists against soil borne pathogens (Kohl et al., 2012; Schmoll, 2010).

Authors in the literature confirmed that molecular approaches led to more refined identification of this genus even at clade and sub-clade level. *Tefl* molecular marker is considered sensitive tool for identification, monitoring and labeling of *Trichoderma* antagonistic candidates during screening procedures. Highly sensitive molecular tools like *tefl* marker are highly efficient barcode for *Trichoderma* species identification regardless confusing morphological characteristics in this genus like color of a colony (Jaklitsch, 2009; Jaklitsch, 2014).

In vitro methods are considered among fast screening techniques and carried out to select *Trichoderma* antagonistic candidates. For instance, *T. asperellum* and *T. harzianum*

have been selected as antagonists against different pathogens using dual culture plates. Dual culture plate has been routinely used to select *Trichoderma* strains with antagonistic activity and omit those who do not present any biological potential (Phillion, 1994; Honor, 1996; Matarese, 2012).

In this study, we attempt to select antagonistic candidates among 17 *Trichoderma* isolates using dual culture assay.

Dual plate test was performed in order to investigate antagonistic potential of *Trichoderma* isolates. Antagonistic activity was evaluated measuring the capacity of *Trichoderma* isolates to inhibit mycelium growth of *Rhizoctonia solani*, *Fusarium oxysporum* and *Verticillium dahlia*; common soil borne pathogens of Oak (Halmschlager and Kowalski, 2004; Matarese et al., 2012).

Translation elongation factor alpha 1 (*tefl*) gene is used as a marker to identify *Trichoderma* isolates at the species level (Jaklitsch, 2011).

MATERIAL AND METHODS

Obtaining *Trichoderma* isolates

Among all fungi isolated from 22 Moroccan soil samples, 17 cultures resembling *Trichoderma spp.* were developed (Table 1).

The strains of *F. oxysporum*, *V. dahlia* and *R. solani* (Table 1) used for the confrontation assays were kindly provided by the Institute of Forest Entomology, Forest Pathology and Forest Protection, University of Natural Resources and Life Sciences, Vienna (BOKU) (IFFF culture collection).

Culture medium

For isolation, cultures were grown on potato dextrose agar medium (PDA, Difco supplemented with ascorbic acid) and were identified by morphological features to genus level.

For genomic DNA extraction and dual culture test, we grew all *Trichoderma spp.* on petri dishes containing Malt extract agar medium which is used to obtain monospore cultures.

Molecular identification of *Trichoderma spp.* by PCR and sequencing of *tefl* DNA

The genomic DNA of *Trichoderma* strains was extracted according to the fast DNA extraction method of fungi (El Khoury, 2011; Schmoll, 2004). The *Trichoderma* strains

Table 1: Fungal cultures used in this study

Cultures	Species	Sites/location	Plants/trees in habitat	Soil types
T2.1; T6.1; T3.2	<i>Trichoderma harzianum</i>	Allal Tazi, Kenitra Morocco	Strawberries culture	Sandy
T4,1	<i>Trichoderma guizhouense</i>	Allal Tazi, Kenitra Morocco	Strawberries culture	Sandy
T9i5; T9i7; T9i8.3; T9i9; T9i10; T9i11; T9i12	<i>Trichoderma reesei</i>	Argania bioreserve, Agadir Morocco	<i>Argania spinosa</i>	Loamy sandy
T9i14; T9i16; T8A1.2; T8A2.3; T8A3.3; T8A4.2	<i>Trichoderma harzianum</i>	Argania bioreserve, Agadir Morocco	<i>Argania spinosa</i>	Loamy sandy
476	<i>Rhizoctonia solani</i>	Patzmannsdorf, lower Austria	<i>Quercus petraea</i>	
539	<i>Verticillium dahliae</i>	Niederweiden, lower Austria	<i>Quercus robur</i>	
145	<i>Fusarium oxysporum</i>	Niederweiden, lower Austria	<i>Quercus robur</i>	

were grown in Malt extract agar at 28°C for 24 hours (hr). *Trichoderma* mycelium was harvested and added to lysis buffer tubes. Then, the samples were incubated at 65°C for 15 minutes (min), allowed to cool in ice and filled with 150 µL of P3 buffer (3.0 M potassium acetate, pH 5.5) containing potassium acetate, glacial acetic acid and sodium hydroxide. After a full speed centrifugation at 15,000 rpm for 10 min, the supernatant was recovered, transferred to new tubes with 700 µL of isopropanol. Full speed centrifugation for 30 min allowed the recovery of the DNA, the supernatant was discarded and the DNA pellet was washed with ethanol and stored at -20 °C in 50 µL of distilled water (El Khoury, 2011; Schmoll, 2004).

Primers were: EF1-728 5'-AGAGTTTGATCCTGGCTCAG-3' and TEF1-LLE 5'-GGTTACCTTGTTACGACTT-3' to amplify the fourth and fifth intron and a part of the large exon of translation elongation factor alpha 1 gene. The amplification reaction was performed in a final volume of 10 µL containing 0.3 µM of each primer, 10 mM each dNTP (Jaklitsch, 2011), 0.1 unit Taq DNA polymerase and 1 µL of DNA (30 ng) and 1 × Taq polymerase buffer. The mixture was first denatured at 95 °C for 2 min, then 40 cycles of PCR (Jaklitsch, 2011) were performed with annealing temperature at 55 °C for 30 seconds (sec) and primer extension at 70 °C for 1 min 30 sec. At the end of the last cycle, the mixture was incubated at 70 °C for 3 min. For each reaction, a negative control without DNA template was included. Efficient amplification was confirmed by gel electrophoresis on 1.5% agarose gel. PCR products were purified and sequencing was done at Eurofins Genomics (Ebersberg, Germany). Phylogenetic analyses were performed in MEGA 6 software program using Neighbor Joining distance algorithm method (Jaklitsch, 2009; Saitou and Nei, 1987; Samuels, 2006). Stability of clades was evaluated by bootstrap rearrangement (1000 replicates) displayed next to the branches (Felsenstein, 1985). Only *Harzianum* clade tree was calculated.

Dual culture test

The *Trichoderma* strains were tested in vitro for their antagonistic activity against *F. oxysporum*, *V. dahliae* and *R. solani* according to the method of Dickinson and Skidmore (1976). The confrontation assays allowed assessing the capacity of the *Trichoderma* strains to inhibit the mycelia growth of the above mentioned phytopathogenic fungi.

Malt extract agar (MEA) discs of 6 mm diameter, cut from the edge of an actively growing colony of each antagonist and pathogen, were placed at opposite sides (4.5 cm from each other) on fresh MEA medium plates (Figure 1a). The radii of the developing pathogen's mycelium were measured in the direction of antagonist's colony (R1) (Figure 1b) three times a day, until contact. Each antagonist/pathogen combination was set up in triplicate and the inoculated plates were incubated at 24±2 °C with a photoperiod of 12 hr/12 hr darkness/light.

The inhibition of mycelia growth in percent PRIG% (Percentage of Radial Inhibition Growth) was calculated after 14 days using the equation (1) ascribed by Skidmore and Dickinson (1976).

$$PRIG = \frac{R_1 - R_2}{R_1} \times 100 \quad (1)$$

Where, R₁: Radius of mycelia growth of pathogenic fungus in control plates (without *Trichoderma*) in mm.

R₂: Radius of mycelia growth of pathogenic fungus in the presence of *Trichoderma* in mm.

Statistical analysis

Data were processed to calculate basic statistical parameters.

The Percentages of Radial Inhibition Growth PRIG% of pathogens mycelia were subjected to one way ANOVA to compare marginal means between different pathogens, two

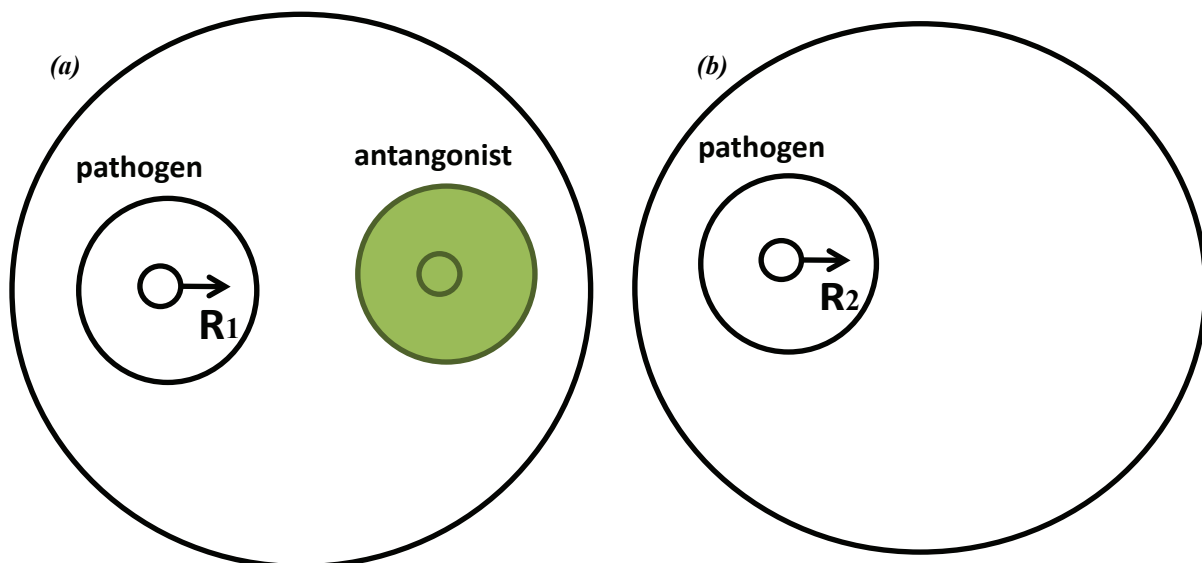


Figure 1: Dual plate culture

(a) dual plates where *Trichoderma* isolate confront pathogen

(b) control plates where pathogen is growing without *Trichoderma*

White disc is positioning of pathogen plug and mycelium and green disc is positioning of *Trichoderma* plug and mycelium

ways ANOVA was conducted to assess interaction effect between pathogens and *Trichoderma* isolates combinations. Statistical analysis of only 14 days data is presented (Korsten, 1993; Korsten, 1995; Skidmore and Dickinson, 1976).

RESULTS AND DISCUSSION

Molecular identification of *Trichoderma*

Trichoderma isolates were identified based on phylogenetic studies of their *tef1* gene. Sequences of *tef1* gene were submitted to NCBI GenBank database. Comparison of *tef1* sequences of isolates to GenBank accessions was performed using nucleotide BLAST alignment program. Sequences alignment revealed similarities with 90 to 99% and 100% homology between *tef1* sequences of isolates and *Trichoderma harzianum* and *Trichoderma reesei* species respectively.

Phylogenetic analysis of *tef1* gene is represented by *Harzianum* clade tree as illustrated in figure 2.

It appears that, *Trichoderma* *tef1* sequences showed variability among isolates and allowed to obtain distinguishably three *Trichoderma* species. Distance methods allowed strong resolution in *tef1* phylogenetic analysis. *tef1* phylogenetic analysis allowed identification and distinction between closely related *Trichoderma* species in the same clade. Therefore, we obtained three different species; *T. afroharzianum* and *T. guizhouense* both belong to *Harzianum* clade (Figure 3) and *reesei* species that belong to *Longibrachiatum* clade.

Previous studies confirmed that *Trichoderma* phylogenetic analysis using *tef1* as molecular marker is more reliable especially for determination of closely related *Trichoderma* species. Elongation factor alpha gene (*tef1*) is highly recommended for the reconstruction of phylogenetic trees and identification of *Trichoderma/Hypocrea*

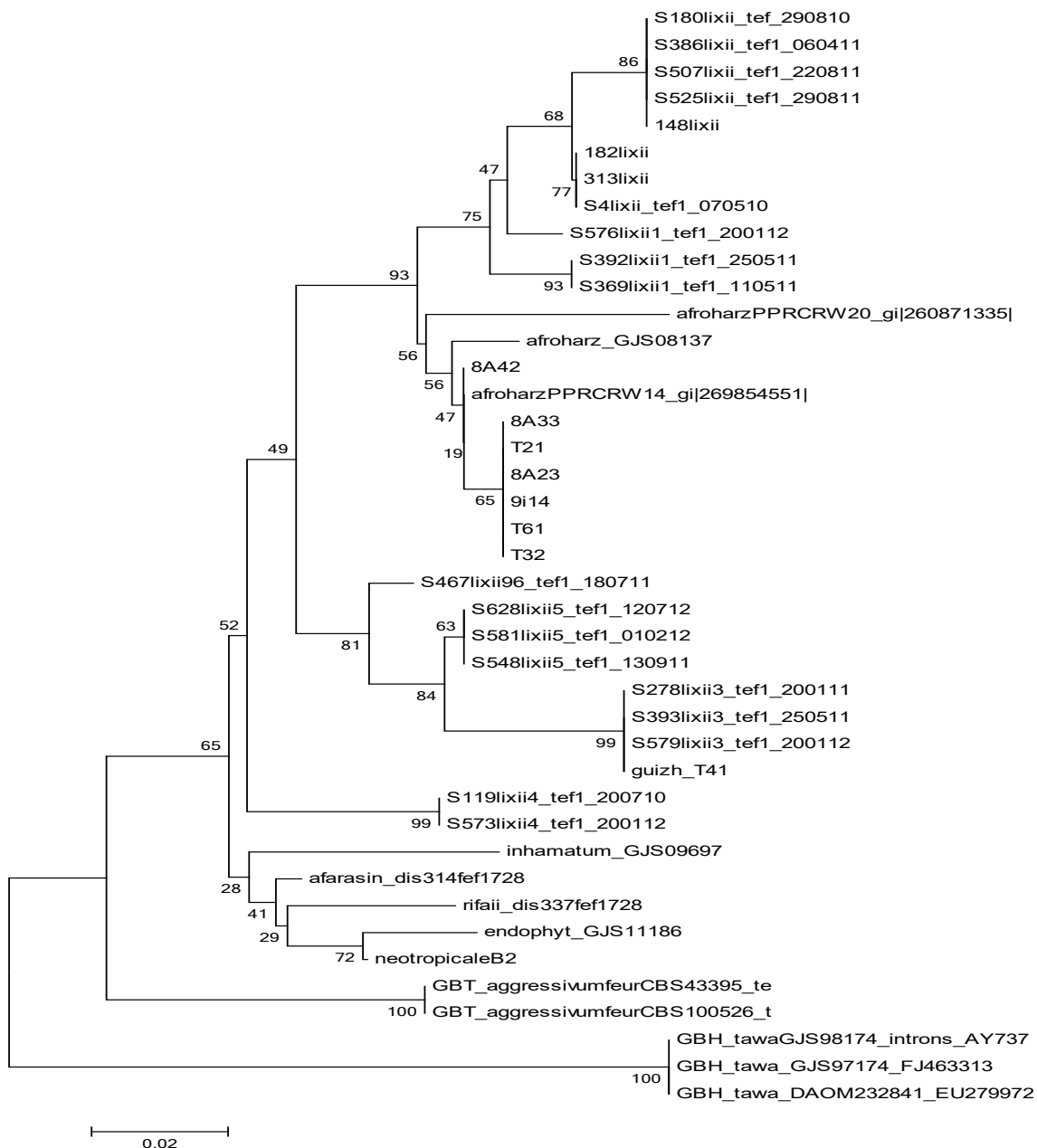


Figure 2: Phylogenetic tree of *Harzianum* clade - The Moroccan strains are attributed to the species *T. afro-harzianum* and *T. guizhouense*. The numbers under branches are bootstrap values.

species to avoid misleading characterization (Kubicek and Druzhini, 2008; Jaklitsch, 2011; Jaklitsch, 2015).

In this study, tef1 phylogenetic analysis allowed allocation of three species according to their habitats distribution (Chaverri et al., 2003; Jaklitsch, 2009; Samuel, 2006).

T. afroharzianum species was isolated from its natural origin in Argania soil in south of Morocco and from Allal tazi soil in North West of the country. However, *Trichoderma reesei* was isolated exclusively from soils of Argania forest in south. T4.1 isolate corresponding to *T. guizouhense* species was sampled from Allal tazi soil in North West of the country. Further isolation and molecular identification must be done for more reliable data interpretation related to geographical distribution of *Trichoderma* species in the country (Jaklitsch 2011; Jaklitsch and Voglmayr, 2015; Kubicek, 2008).

Antagonistic activity of *Trichoderma* isolates

Dual plate culture method showed considerable antagonistic activity against the three different pathogens when confronting 17 *Trichoderma* isolates and results of PRIG% means are presented in Table 2.

The Tukey range test ($\alpha=0.05$) assorted nine distinguishing groups of *Trichoderma* isolates depending on different PRIG% intervals. First group contains the isolate 8A2.3 with 95.3% as the highest PRIG% mean against three pathogens. Second group contains T3.2 and 8A3.3

reducing the radial mycelia of pathogens by 94.0%, third group contains 8A4.2, 8A1.2, T4.1 and T9i12 by 93.0%, the fourth group T6.1, T9i16, T9i10 by 89.0% - 83.0% PRIG% intervals followed by fifth group T9i9 and T9i8.3 by 82.0% and the sixth group T9i7, T9i11 reducing radial growth of pathogens mycelia by 81.0% - 80.0% respectively. Other orphan groups containing one isolate each are cited as following; T2.1 with 89.9%, group T9i5 with 81.9% and group T9i14 with 77.3%.

Korsten (1995) developed a growth inhibition categories on a scale ranking antagonistic in vitro screening by dual plate culture where 0 scale = no inhibition growth, 1 scale = 1 to 25% inhibition, 2 scale = 25% to 50%, 3 scale = 50% to 75% and 4 scale = 75% to 100%. Therefore, we proposed to scale our *Trichoderma* isolates using Korsten antagonism scale in Table 2.

Almost all *T. afro-harzianum* isolates T8A.3.3, T8A.2.3, T8A.4.2, T8A.1, T6.1, T3.2, T9i14.1, T2.1 et T4.1 showed the highest inhibition percentage compared to the other *T. longibrachiatum* isolates T9i5, T9i8.3, T9i7, T9i9, T9i10, T9i11, T9i16, with less antagonist performance against the three pathogens with exceptions like T9i12 as shown in Table 2.

Analysis of variance showed differences in inhibition activity assessed by a significant PRIG% means with $P = 0.000$ ranking *F. oxysporum* with highest PRIG% means followed by *V. dahlia*, and by *R. solani*.

Table 2: Inhibition of the growth of *Rhizoctonia solani*, *Verticillium dahlia* and *Fusarium oxysporum* by 17 *Trichoderma* isolates and their antagonistic activity against those pathogens in dual culture test

<i>Trichoderma</i> isolates	<i>Rhizoctonia solani</i>		<i>Verticillium dahlia</i>		<i>Fusarium oxysporum</i>	
	PRIG% means (%)	Antagonistic activity (on 1-4 scale)	PRIG% means (%)	Antagonistic activity (on 1-4 scale)	PRIG% means (%)	Antagonistic activity (on 1-4 scale)*
<i>Trichoderma afro-harzianum</i>						
8A1.2	96.6	4	90.7	4	93.3	4
8A2.3	98.3	4	93.2	4	94.4	4
8A3.3	95.8	4	92.7	4	93.9	4
8A4.2	96.6	4	89.7	4	94.4	4
T2.1	88.9	4	85.8	4	95.0	4
T3.2	99.1	4	93.6	4	89.7	4
T6.1	89.8	4	91.2	4	86.9	4
T9i14	83.9	4	76.6	4	71.7	3
<i>Trichoderma guizouhense</i>						
T4.1	100	4	93.6	4	86.9	4
<i>Trichoderma longibrachiatum (reesei)</i>						
T9i10	99.1	4	84.4	4	67.2	3
T9i11	91.5	4	80.5	4	69.4	3
T9i12	97.4	4	89.3	4	93.3	4
T9i16	95.8	4	89.7	4	72.8	3
T9i5	93.2	4	85.4	4	67.2	3
T9i7	95.8	4	80.5	4	67.2	3
T9i8.3	95.8	4	78.0	4	73.6	3
T9i9	91.5	4	79.0	4	76.9	4

* Korsten antagonism scale (1995)

Where 0 scale = no inhibition growth, 1 scale = 1 to 25% inhibition, 2 scale = 25% to 50%, 3 scale = 50% to 75% and 4 scale = 75% to 100%. All PRIG% data represent means of three independent experiments +/- Standard deviation

Two ways ANOVA variance analyses show significant interaction $P = 0.000$ in *Trichoderma* isolates – pathogen combinations. PRIG% values were statistically significant ($P \leq 0.05$) for each *Trichoderma* isolate confronting different pathogens suggesting that radial growth inhibition of pathogen mycelium depends tightly on the presence of the antagonist *Trichoderma* (Rahman et al., 2009).

Antagonistic activity was also evaluated by observation of *Trichoderma* overgrowth up on pathogens and spores production and proliferation in dual plates.

It seems that all *Trichoderma* colonies overgrow up on pathogens colonies and that was visible after few days of both fungi contact in dual plate. *Trichoderma* isolates produced spores when confronting pathogens and proliferate once their mycelium cover pathogens mycelium. No color changes have been noticed in *Trichoderma* cultures (Prisa, 2011).

DISCUSSION

Prior work has documented the effectiveness of many *Trichoderma* species as biocontrol agents to control plant pathogens thanks to their combined antagonistic mechanisms. For instance, Lorito and co-workers reported that antibiosis is one of the mechanisms of *Trichoderma* antagonistic candidates. They focused on effect of secondary metabolites and antibiotics production on pathogens growth cease. Other studies focused on mycoparasitic mechanism based on production of degrading enzymes (i.e. chitinases and β -1,3-glucanases) (Chet, 1987; Lorito et al., 2010).

Simple in vitro screening techniques like dual culture plate are used to select varieties of antagonistic profiles of the genus. Dual plate technique is widely used to discover antagonistic potential of *Trichoderma* spp. and its direct interaction with the pathogen.

Another point, researchers have demonstrated high sensitivity of molecular marker to detect, identify and monitor the fungus of interest during screening procedures. Jaklitsch have recently worked on advanced molecular strategies for reconstruction of *Trichoderma* clades. Jaklitsch and co-workers have demonstrated that ITS, *tefl* and *rpb2* are remarkably suitable for *Trichoderma* identification. They reported that using *tefl* alone seems to be satisfactory to discriminate between closely related species of this genus. In the other hand, integrated genetic analysis is crucial for specific characterization of the antagonistic profiles of different species which is a costly process (Jaklitsch, 2006; Jaklitsch, 2011).

Therefore, in this study we tested the effectiveness of *tefl* phylogenetic study in identifying *Trichoderma* isolates. This study compares well with previous *tefl* phylogenetic analysis. Furthermore, we attempt to establish a preliminary screening of antagonistic candidates among 17 *Trichoderma* isolates.

We found that in vitro test resulted in labeling some antagonistic candidates among *Trichoderma* isolates based on dual culture test. That is, significant high inhibition of pathogens' mycelium was recorded. Moreover, all 17 *Trichoderma* isolates performed tropic growth over almost all pathogens mycelium.

It can be inferred that dual culture test is promised selection method of antagonistic candidates during screening procedure. However, it must be applied with cautions. In fact, dual plate is a small esthetic micro-environment that does not show the real behavior of *Trichoderma* isolates. Therefore, further green houses and field experiments are highly recommended to evaluate antagonistic activity implicating other components.

With *tefl* analysis we could label antagonistic candidates among *Trichoderma* spp. by identifying isolates at the species level. Therefore, *Trichoderma harzianum* species isolated from Argan forest (i.e. T8A4) were able to inhibit significantly growth of pathogens. However, *T. reseei* isolates showed less promising antagonistic effect during dual confrontation test except T9i12 *reseei* isolate. T9i12 inhibits significantly pathogens' growth.

Other observations for antagonistic evaluation were overgrowth and sporulation over pathogens colony. *Trichoderma* spp. showed interesting antagonistic interaction like over growth and sporulation over three pathogens.

This finding encourages for more tests based on microscopic observation of short loops, coiling and degradation of pathogens mycelium when in contact with *Trichoderma* hyphae.

Most notably, *Trichoderma* spp. inhabiting Moroccan soil; *Argania* forests, where climate conditions are semi-arid, represent so far an overlooked habitat to isolate *Trichoderma* spp. Therefore, identifying and classifying *Trichoderma* species inhabiting new ecological niches like *Argania* forests may lead us to detect interesting antagonistic candidates (Khattabi, 2004; Druzhinina et al., 2010).

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