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Biological insect control using Metarhizium anisopliae: morphological, molecular, and ecological aspects
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ABSTRACT

Microbial control of insects is based on the rational use of pathogens to maintain environmentally balanced pest population levels, and Metarhizium anisopliae has been the most studied and most utilized fungal species for that purpose. The natural genetic variability of entomopathogenic fungi is considered one of the principal advantages of microbial insect control. The inter- and intraspecific variability and the genetic diversity and population structures of Metarhizium and other entomopathogenic fungi have been examined using ITS-RFLP, ISSR, and ISSP molecular markers. The persistence of M. anisopliae in the soil and its possible effects on the structures of resident microbial communities must be considered when selecting isolates for biological insect control.

Key words: biological control, Metarhizium anisopliae, molecular markers.

INTRODUCTION

Biological control consists of the introduction of beneficial predatory or parasitic species into cultivation systems where they were previously absent or present only at low population levels. This technique is designed to negatively affect specific target species that could otherwise become pests or infectious agents (GLIESSMAN, 2001). Susceptibility to pests is a general reflection of plant health, which can be negatively influenced by poor soil fertility management (NICHOLLS & ALTIERI, 2007). One of the objectives of biological control is to assure that the beneficial organism to be introduced can complete its lifecycle at the site, and then reproduce with sufficient efficiency to become a permanent resident of the agrosystem. Frequently, however, the niche conditions available to the beneficial introduced organism do not fully satisfy its long-term needs, requiring its reintroduction (GLIESSMAN, 2001). Changes in production practices and the use of agricultural additives are often necessary for biological control to be successful. Integrated Pest Management (IPM) is an alternative to unilateral intervention strategies using agrochemicals, with a wider focus on the ecology of the insect pests as well as the crop plants, based on the use of complementary tactics and the adoption of cultivation techniques that favor plant diversity. Pest control in this type of approach is initially based on natural agents such as pathogens,
parasites and predators, with the use of agrotoxins being contemplated only as a last resort. However, as biological pest control methods do not demonstrate immediate results in agro-industrial systems with large-scale production and commercialization goals (as agrotoxins), commercial groups tend to avoid the costs and labor related to their development and perfection. Nonetheless, growing energy costs, environmental degradation, and inflation all reinforce the argument that immediate financial gains should not be the principal motivating force in agricultural production (ALTIERI, 2002). In spite of the strong economic pressure on agricultural production, many farmers are making the transition to practices that are more environmentally friendly and have the potential to contribute to long-term agricultural sustainability with biological control being one of the principal tools in this conversion process (GLIESSMAN, 2001).

Microbial control is an aspect of biological insect control and consists of the rational use of pathogens to maintain pest balances in agricultural environments, with increases in the numbers of other natural enemies often being observed in fields where microbial control has been used. Successful programs of microbial control using entomopathogenic fungi to combat arthropod pests in soils and aquatic environments have been developed, principally utilizing the genera Metarhizium, Beauveria, Sporothrix, Lecanicillium, Nomuraea, Hirsutella, Aschersonia, Isaria, Paecilomyces, and Enthomophthora (ALVES & LOPES, 2008). Species within the genus Metarhizium are pathogenic fungi having broad ranges of insect hosts. M. anisopliae was found to be a species complex composed of nine species based on multilocus phylogeny (BISCHOFF et al., 2009). The objective of this study was to analyze some morphological, molecular and ecological aspects of M. anisopliae.

Metarhizium anisopliae

*Metarhizium anisopliae,* a anamorphic fungus which belong to the phylum Ascomycota, is the most intensively studied species of the genus *Metarhizium,* considering that the teleomorph *Cordyceps brittlebankisoides* [= *Metacordyceps brittlebankisoides* (Liu, Liang, Whalley, Yao & Liu)] Sung, Sung, Hywel-Jones & Spatafora] was isolated from insect larva (*Coleoptera: Scarabaeidae*) and identified as *M. anisopliae* var. *majus* [= *M. majus* (Johnston) Bischoff, Rehner & Humber] (LIU et al., 2001). The reproductive structures of *M. anisopliae* (the anamorph, the most commonly encountered form) comprise conidiophores and conidia. Leveduriform structures or blastospores and appressoria are produced by *M. anisopliae* through mycelial differentiation. Blastospores can function in certain cases as reproductive units and are produced in submerged cultures (JACKSON & JARONSKI, 2009) and in the hemolymph of insect hosts (ALVES, 1998). The appressoria, formed at the extremity of the hyphae, may be involved in fungus pathogenicity and have the function of initiating epicuticular and prouticule penetration of the insect tegument (ALVES, 1998). The production of microsclerotia by isolates of *M. anisopliae* has been observed after cultivation in liquid media with different concentrations of carbon and carbon-nitrogen (JACKSON & JARONSKI, 2009).

The fungal-host relationship occurs through the adhesion and germination of conidia on the surface of the insect, followed by hyphae penetration through the cuticle. The process of host colonization initiates after penetration, with the penetrating hyphae becoming thicker and ramify within the tegument and the hemocoel of the insect, forming blastospores. The hyphae continue to grow and invade various internal organs after the death of the host and will subsequently emerge from the insect body and produce conidia that disseminate and infect other individuals (ALVES, 1998).

Molecular studies of the processes of host infection have shown them to be complex and multifactorial. The adhesion and penetration steps have been most closely examined and appear to be decisive to infection. The participation of an adhesin coded by the gene *Mad1* in the adhesion of conidia to the cuticle of *Manduca sexta* Linnaeus larva was demonstrated using mutants in which this gene was deleted, with these mutants demonstrating significant decreases in conidial germination, suppression of the formation of blastospores, and reduced virulence (WANG & St. LEGER, 2007a). COSENTINO-GOMES et al. (2013) described that the inhibition of phosphatase activity in the conidia of *M. anisopliae* reduced adhesion to the integument of *Dysdercus peruvianus* (Hemiptera: *Pyrrhocoridae*) and (indirectly) its infection.

The participation of perilipin (proteins that surround lipidic droplets in the cell interior) in appressoria differentiation in *M. anisopliae* has also been reported. The deactivation of the *Mpl1* gene in some strains generates deficiencies in the infection process due to the formation of appressoria with lower concentrations of lipidic droplets and resultantly lower levels of osmotic pressure - resulting in difficulties in terms of hyphal penetration (WANG, 2002).
Defective appressoria were also observed after the deletion of the \textit{mapk1} gene (catalytic subunit 1 of the protein kinase A) (FANG et al., 2009). Subtilisin-type proteases have been intensively studied in penetration processes, and 10 genes are known to code for different isoforms of these enzymes (Pr1A - Pr1J) and appear to reflect specificity in relation to different hosts (BAGGA et al., 2004). MOS1 is another protein with an apparent role in the adaptation of fungi to the high osmotic pressure encountered in insect hemolymph (WANG et al., 2008). Other genes, such as \textit{Mcl1} (collagen-like protein), \textit{Cog8} (which regulates the G protein signaling pathway), \textit{chi2} (endochitinase), \textit{chi3} (endo- and exochitinase), and \textit{Mpk1} (phosphoketolase) are known to be involved in the host infection processes of \textit{M. anisopliae}, with reductions in virulence if they are inactivated (FANG et al., 2007; BOLDO et al., 2009; DUAN et al., 2009). SU et al. (2013) undertook comparative proteomic analyses of the conidia and mycelia of \textit{M. anisopliae} (Ma1291). The proteins identified as exclusive to the conidia were involved in protective processes, appressorium formation, and the degradation of the host cuticle and exclusive proteins to mycelia were involved in biosynthetic and energy-generating metabolic processes, such as UTP-glucose-1-phosphate uridyltransferase and heat-shock protein 70.

Molecular characterization

Molecular markers can represent the phenotype of an expressed gene or a DNA segment corresponding to a non-expressed region of the genome. Advances in molecular biology have resulted in the development of various methods for detecting genetic polymorphism at the DNA level and have aided our understanding of genetic diversity and the population structures of fungi populations (FALEIRO, 2007).

The polymerase chain reaction (PCR) technique, allied to methodologies of cloning and DNA sequencing, have allowed the rapid accumulation of information relating to genome structure and the discovery of repetitive DNA sequences (which are rich sources of genetic polymorphism). A number of methodologies have been described for analyzing polymorphisms based on PCR, including ITS-RFLP (Internal Transcribed Spacer - Restriction Fragment Length Polymorphism), ISSP (Intron Splice Site Primer), ISSR (Inter Simple Sequence Repeats), and SSR (Simple Sequence repeats) (FALEIRO, 2007).

The DNAs coding for rRNA are arranged as genetic aggregates with three genetically conserved regions composed of 18S, 5.8S and 28S genes that are transcribed and processed to generate mature RNA, but are separated by variable intergenic spacer regions denominated ITS1 and ITS2. The genetic aggregate that codes for rRNA appears to be repeated hundreds of times in the fungus genome and demonstrates both highly conserved and variable regions, allowing scientists to analyze variations at different taxonomic levels. The 18S region is the most highly conserved, and is therefore only used in comparisons between distantly related organisms. The 28S region is more variable and therefore appropriate for comparing different genera (or different species, in some cases). ITS regions evolve relatively rapidly and can be used to distinguish closely related species or even varieties within the same species (FUNGARO, 2000).

DNA samples digested with restriction enzymes (RFLP) can identify polymorphisms based on the numbers and sizes of the fragments produced, which allows the differentiation of species and isolates of \textit{Metarhizium} based on the presence or absence of rDNA restriction sites (PIPE et al., 1995). In the present study, isolates of \textit{M. anisopliae} could be grouped according to their geographical origins, although no significant correlations were observed in terms of their hosts. VELÁSQUEZ et al. (2007) observed that there were no associations between the diversity of isolates of \textit{M. anisopliae} from different regions in Chile and their geographic origins. ITS-RFLP was used to define specific primers that could be used to detect and identify \textit{M. anisopliae} \textit{var. anisopliae} (DESTÉFANO et al., 2004).

Eukaryotic chromosomes contain genes that are separated by non-coding regions (introns) as well as regions with coding information represented by proteins (exons). Introns can be separated into four basic categories according to their structural characteristics and self-splicing mechanisms: group I, II, nuclear pre-mRNA, and nuclear tRNA. The introns of groups I and II are classified according to their internal organizations and have the intrinsic capacity of self-splicing; the latter two intron groups can be used as molecular markers in intra- and interspecific studies of diversity (HAUGEN et al., 2005).

Group I introns are encountered in eukaryotic organisms such as fungi, protists, and green algae in nuclear, mitochondrial, and chloroplast genomes. Group I introns are encountered in the eukaryotic nuclear genome in rDNA genes at specific sites that code for the larger and smaller rRNA subunits. These introns are autonomous genetic elements characterized by their capacity to transfer from one allele to another (as some are mobile.
elements – transposons) and by their ability to self-spool from RNA transcripts (Haugen et al., 2005). Group I introns are generally irregularly distributed – being present in some isolates but absent in others – and thus can serve as markers of genetic variability. Genetic diversity among Metarhizium anisopliae isolates have been observed in studies of genes associated with the largest ribosomal subunit and with four insertion sites of group I introns, and the presence/absence of these introns allow the delimitation of seven groups (Marquez et al., 2006).

Microsatellite DNAs show numerous short, repeated, tandem sequences, and their analyses involve replicating fragments containing those repetitions through the use of oligonucleotides that bind to the regions which flank them (SSR sites). These markers were used to examine polymorphism in Metarhizium anisopliae, but the primers used to examine samples derived from soil cores from different regions in Chile (Velásquez et al., 2007) and numerous countries in Asia and Europe (Freed et al., 2010) were not efficient in detecting informative polymorphisms. To the contrary of microsatellite analysis, the ISSR technique amplifies fragments located between two repetitive regions present in various genomes (Faleiro 2007), known as inter-microsatellite regions. This ISSR marker did demonstrate differences among different isolates of Metarhizium anisopliae var. anisopliae of the same origin and from the same host, principally when using the primers (GACA)$_4$ and (GTG)$_5$, providing DNA fingerprints for a number of isolates (Tiago et al., 2011). The genetic structure of Metarhizium spp. (Metarhizium anisopliae and its sister species, M. robertsi), pathogens found in Chinese burrower bugs populations (Schiodetta formosana), were assessed using ISSR. They differentiated into two main clades including over 71% of all strains causing epizootics, with a similarity of 83% (Luan et al., 2013).

The soil ecology of Metarhizium anisopliae

Metarhizium anisopliae demonstrates considerable metabolic and ecological versatility and has been observed colonizing the rhizosphere and adhering to the surfaces of plant roots, and it may significantly influence this ecological niche by repelling and killing soil insects (Hu & St. Leger, 2002). Bruck (2005) observed that the conidia of Metarhizium anisopliae demonstrated greater persistence in the rhizosphere of Picea abies Linnaeus than in the soil alone. On the other hand, a pilot study by St. Leger (2008) in a pasture site indicated that Metarhizium anisopliae could survive for various years in the soil, although with fluctuations in its population levels. Additional studies will therefore be necessary to determine if the plant rhizosphere can truly be considered a refuge (a locality where the fungus can survive outside its insect host) for Metarhizium anisopliae in the soil. Meyling & Eilenberg (2007) suggested that plant associations are important to the biological cycle of Metarhizium anisopliae in temperate regions. It is possible that this fungus has multiple functions in terms of plant protection, with antagonistic effects against phytopathogenic fungi.

A number of studies have examined the molecular mechanisms involved in the capacity of Metarhizium anisopliae to adhere to both insects and roots, resulting in the identification of adhesins MAD1 and MAD2. Adhesin MAD1 is involved in insect pathogenicity and MAD2 with fungal adhesion to plant roots (Wang & St. Leger, 2007a), and a study by Wang et al. (2005) examining genetic expression demonstrated that Metarhizium anisopliae could act as both a pathogen (growing on the cuticle and in the hemolymph of insect hosts) and a saprophyte in the rhizosphere (growing on the bean root exudates). Wyrebek & Bidochka (2013) amplified and cloned the full Mad1 and Mad2 genes in fourteen isolates of seven different species of Metarhizium to assess their genetic variability. Phylogenetic analyses of 5' EF-1α (which is used for species identification), Mad1, and Mad2 indicated that the evolution of the Mad2 gene was more congruent with the phylogeny of 5' EF-1α than of Mad1. This suggests that Mad2 diverged among the Metarhizium lineages and contributed to clade- and species-specific variations, while Mad1 was largely conserved.

Studies have shown that some insect pathogenic endophytic fungi, such as Metarhizium, are able to transfer insect-derived nitrogen to plant roots, probably in exchange for plant sugars. Metarhizium has a phylogenetic heritage of plant symbiosis (i.e., the genus is closely related to other endophytes) and has also evolved as a generalist insect pathogen (Behie et al 2013).

A number of workers have investigated the persistence of Metarhizium anisopliae in the soil, with greater fungal survival being observed in sandy-clay soils and in soils with average compaction density values (Lanza et al., 2004). High average numbers of colonies of Metarhizium anisopliae could be recovered 30 days after inoculation, and viability for up to 120 days was observed in previously sterilized soils (Guerra et al., 2009) and for up to 216 days after field inoculation (Martins et al., 2004). A study based on quantitative PCR (qPCR) demonstrated that
**Metarhizium** Clade 1 (*M. majus*, *M. guizhouense*, *M. anisopliae*, *M. robertsii*, *M. brunneum*) was present at high densities in soil samples from pastures and improved brunneum *M. pinghaense* in insecticide Carbofuran, which demonstrated significant differences in the soil fungal community; this result differed from the situation after using the transitory effects on the soil fungal community; without the presence of its insect host had only small activities occurring in that milieu. The introduction of exogenous microorganisms into natural and agricultural ecosystems may affect the soil microbial community and, consequently, diverse ecological processes in those environments. The effects of the introduction of entomopathogenic fungi into soil microbial communities represent an ecological intervention that has not yet been extensively examined. A study by SCHWARZENBACH et al. (2009) using ribosomal internal spacer analysis (RISA) to examine the effects of *B. brongniartii* on fungal community structures in soil microcosms indicated that its presence in the soil microcosms indicated that its presence in the soil only small (or transitory) effects on the soil fungal community; this result differed from the situation after using the insecticide Carbofuran, which demonstrated significant impacts even at the end of the experimental period. Other studies using the DGGE technique indicated that the fungal soil community structure was not significantly influenced by the presence of *M. anisopliae* var. *anisopliae* URM5951 at 15, 30, 60, and 90 days after its inoculation in the soil (TIAGO et al., 2012).

**CONCLUSION**

Molecular genetic techniques can be useful in addressing taxonomic problems and in determining degrees of inter- and intraspecific genetic variation. Polymorphism studies have contributed to our understanding of the genetic diversity and population structures of fungi and have provided information that can be very important to programs of biological control. Molecular biological techniques have important roles in our understanding of the genes involved in host infection processes, such as the adhesion steps, appressorium formation, and the degradation of the host cuticle. Measures of the persistence of entomopathogenic fungi in the soil and any possible effects of their application on the structural and functional diversity of soil or rhizosphere microbial communities are essential ecological aspects that must be understood in agrosystems, as these microorganisms represent a significant fraction of the soil biota in terms of its species diversity and the multiplicity of metabolic activities occurring in that milieu. *Metarhizium* is a plant symbiont that can act as a saprophyte in the rhizosphere but has also evolved as a generalist insect pathogen. As such, the paradigm that *M. anisopliae* is principally an insect pathogen is questionable, and additional studies will be necessary to better understand its ecological role in the soil.

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