



## Molecular Identification of Etiological Agents of Chromoblastomycosis in Costa Rica

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### Abstract

Chromoblastomycosis is the second most frequently reported subcutaneous mycosis in Costa Rica. It is caused by dematiaceous fungi belonging to the family Herpotrichiellaceae (Order Chaetothyriales), especially by *Fonsecaea pedrosoi* and *Cladophialophora carrionii*. However, it is important to note that *Fonsecaea monophora* is able to disseminate and cause cerebral phaeohyphomycosis. Thus, five clinical isolates deposited in the Fungal Collection of the School of Microbiology of the University of Costa Rica were analyzed. The isolates were characterized macroscopically and microscopically after grown in potato dextrose agar. Genetic identification was performed via amplification and sequencing of the ITS (internal transcription spacer) region. The isolates were identified as *F. pedrosoi* (n = 3), *F. monophora* (n = 1) and *Rhinochadiella aquaspersa* (n = 1). Hence, we report for the first time that *R. aquaspersa* is an etiological agent of chromoblastomycosis in Costa Rica and confirm the presence of both *F. pedrosoi* and *F. monophora* in the country. Therefore, we recommend the usage of molecular techniques to identify these pathogens since there is a risk of fungal dissemination in our patients.

**Keywords:** Chromoblastomycosis; Chromomycosis; *Fonsecaea monophora*; *Fonsecaea pedrosoi*; *Rhinochadiella aquaspersa*; Subcutaneous Infections

### Abbreviation

ITS: internal transcription spacer

### Introduction

Chromoblastomycosis is a worldwide disease, however, most cases are reported in tropical and subtropical areas, especially in Latin America [1-5]. It is the second most frequently reported subcutaneous mycosis in Costa Rica, surpassed only by sporotrichosis [6,7]. Thus, in 1953, Costa Rica ranked third in incidence (n = 53 cases), below Brazil and Cuba [2].

It is a cutaneous and subcutaneous infection that is obtained by traumatic inoculation of organic material contaminated with

dematiaceous fungi. It usually affects lower limbs and is slowly evolving as the infection grows approximately one centimeter per year. A papule or plaque appears at the site of inoculation, which then acquires a verrucous form. Older lesions can ulcerate, become infected secondarily, and eventually culminate in squamous cell carcinoma. Also, lesions in the form of atrophic plaque have been described. In direct examinations (potassium hydroxide or histological stains) Medlar bodies are observed [1-5].

The etiologic agents are from the Herpotrichiellaceae family (order Chaetothyriales) among which are the species *Fonsecaea* spp., *Cladophialophora carrionii*, *Phialophora verrucosa*, *Exophiala jeanselmei* and *Rhinochadiella aquaspersa* [8-15]. In regard to the

genus *Fonsecaea*, the usage of molecular biology has discerned that it is composed of several species, being those of medical importance: *F. pedrosoi*, *F. monophora* and *F. nubica* [16]. This is of great importance because at the microscopic level these fungi are very similar, however, the pathology they cause differs. *F. pedrosoi* and *F. nubica* are exclusive causative agents of chromoblastomycosis, while *F. monophora* can cause other clinical conditions such as cerebral phaeohyphomycosis [17-19] and/or spread to lymph nodes, lungs, kidneys and liver [17]. Hence, the diversity and complexity of the clinical manifestations caused by *F. monophora* is the reason why various groups of researchers, worldwide, have been given the task of characterizing at the molecular level all those isolates previously identified as *F. pedrosoi*, by light microscopy techniques. For example, Yaguchi *et al.* (2007) [17], analyzed 20 Japanese strains of which 100% were identified as *F. monophora* and 21 Latin American strains, of which 47.62% ( $n = 10$ ) were identified as *F. pedrosoi* and 33.33% ( $n = 7$ ) as *F. monophora*. Similar results were published by de Hoog, *et al.* (2004) [10] and by Najafzadeh, *et al.* (2009) [11].

To date, no work has been carried out in Costa Rica that continue with these international initiatives. This is essential since the study of Yaguchi *et al.* (2007) [17] analyzed three Costa Rican clinical isolates out of which one was molecularly identified as *F. monophora*, thus proving the existence of this fungus in our country. In turn, it is very important to know which are the causative agents of this pathology and to evaluate whether these fungi can also be potential agents of invasive phaeohyphomycosis. Therefore, the present study aims to identify by molecular biology techniques the isolated chromoblastomycosis causing fungi from the collection of the Medical Mycology Laboratory of the School of Microbiology, University of Costa Rica.

## Materials and Methods

### Clinical isolates

Five Costa Rican clinical isolates, from patients suffering chromoblastomycosis were analyzed; these isolates were deposited in the Fungal Collection of the Section of Medical Mycology of the School of Microbiology, University of Costa Rica. The fungi were kept in potato dextrose agar (PDA), with a layer of mineral oil and at room temperature (20 - 30°C). *Fonsecaea pedrosoi* CBS 659.76 and *F. monophora* CBS 269.37 were used as control strains.

### Phenotypic characterization

A drop of clear lactophenol reagent was placed on a slide and then a small portion of the fungal colony was placed on top of it, then, the preparation was covered with a coverslip and slightly

heated in the flame, without boiling. Finally, the preparation was observed under a microscope (400x) so that the typical morphology of each fungus could be observed [20].

### DNA extraction and amplification

DNA extraction was performed using The Nucleo Spin® Tissue kit (Macherey-Nagel, Germany), following manufacturer procedure. The internal transcription spacer (ITS) was amplified according to the protocol described by White, *et al.* (1990) [21], using the following pair of primers: ITS1 (5'-TCC GTA GGT GAA CCT GCG G-3') [forward] and ITS4 (5'-TCC TCC GCT TAT TGA TAT GC-3') [reverse]. The PCR products were sent to Macrogen Inc. (South Korea), where the sequencing process was carried out. The sequences obtained were reviewed and edited using the BioEdit v 7.0.5.2 [22] and aligned with reference sequences from the NCBI database in MEGA [23]. Multiple sequence alignments were performed with ClustalX [24]. Phylogenetic trees were constructed using Bayesian Inference through the MrBayes 3.2.6 program; 1000 replicates were analyzed; the resampling percentage was reflected in each branch of the tree.

## Results and Discussion

### Clinical isolates and phenotypic characterization

Chromoblastomycosis is the second most frequently reported subcutaneous mycosis in Costa Rica, preceded only by sporotrichosis. As for its epidemiology, most cases occur in adult male patients and the lesions are located in the lower limbs [14]. The isolates analyzed in this study mostly come from male patients, but it is important to highlight that the majority of the patients had lesions in their upper limbs; situation which, although not common, has also been reported in the literature [12,25,26]. Demographic data of the patients is found in table 1. Only the isolate FOPE 26 had information of the year in which it was deposited in the fungal collection: 2002. Macroscopic morphology of the fungal cultures was composed of black and velvety texture colonies. At the microscopic level, the isolates FOPE 02, 17, 22 and 26 presented dematiaceous, septate mycelium with the presence of branched conidiophores, simple acroteca-type conidiophores and/or phyalid-type conidiophores; meanwhile, FOPE 04 presented dematiaceous, septate mycelium with the presence of only simple acroteca-type conidiophores (Figure 1).

### Molecular identification

Prior to this work, Costa Rican studies focused only on the morphological identification of the etiological agents of chromoblastomycosis [8,13,14], where those fungi that presented dematiaceous,

Isolate	Preliminary identification*	GenBank access code	Patient Demographics	
			Gender	Anatomical site of the lesion
FOPE 02	<i>Fonsecaea</i> sp.	MH374865	Female	Arm
FOPE 04	<i>Rhinocladiella</i> sp.	MH374866	NI	Arm (elbow)
FOPE 17	<i>Fonsecaea</i> sp.	MH374872	Male	Little finger
FOPE 22	<i>Fonsecaea</i> sp.	MH374867	Male	Right ankle
FOPE 26	<i>Fonsecaea</i> sp.	MH374868	Male	Leg

**Table 1:** Preliminary identification of the isolates analyzed and demographic data of patients with chromoblastomycosis from which they were obtained.

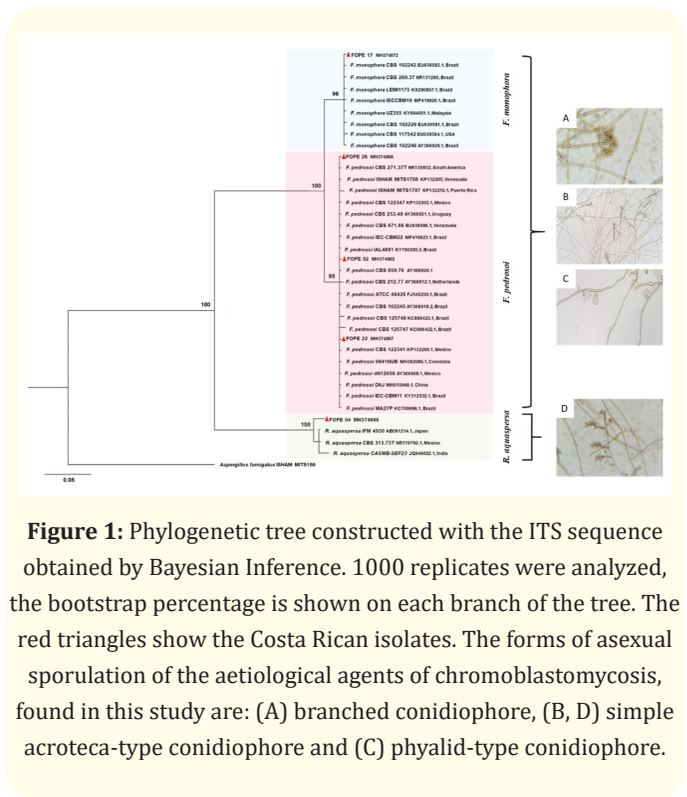
\*: Microscopic identification of the saprophytic phase.

NI: Not Indicated.

septated mycelium with the presence of two or three of the following types of asexual sporulation: branched conidiophores, simple acroteca-type conidiophores and simple phyalid-type conidiophores were classified as members of the genus *Fonsecaea*. However, in 2004, de Hoog, *et al.* [10] studied *Fonsecaea* isolates based on the analysis of their ITS sequences and random amplification of polymorphic DNA (RAPD), finding a high diversity and proposing the separation of the genus in two species: *F. pedrosoi* and *F. monophora*. Over the years, new studies have defined two more species: *F. nubica* and *Fonsecaea multimorphosa*.

Species-level identification is related to the pathology they cause, with *F. pedrosoi* and *F. nubica* been exclusive agents of chromoblastomycosis, and *F. monophora* and *F. multimorphosa* causing cerebral phaeohyphomycosis in immunocompetent patients and felines, respectively [17,19,27]. On the other hand, fungal infections at the brain level caused by hyaline fungi such as *Cryptococcus* spp., *Aspergillus* spp., *Fusarium* and *Mucorales* are mostly associated with immunosuppressive patients [28-30].

The sequences obtained were deposited in GenBank. The access codes are shown in table 1. The phylogenetic analysis was performed using type strain sequences in order to estimate with greater certainty the taxonomy of the clinical isolates. The analysis grouped the isolates sequences under study into three groups: n = 3 were identified as *F. pedrosoi*, n = 1 as *F. monophora* and n = 1 as *R. aquaspersa* (Figure 1). The evolutionary history was inferred using Bayesian Inference. The consistency index is (0.953271), the retention index is (0.985549), and the composite index is 0.959884 (0.939495) for all sites and parsimony-informative sites (in paren-



**Figure 1:** Phylogenetic tree constructed with the ITS sequence obtained by Bayesian Inference. 1000 replicates were analyzed, the bootstrap percentage is shown on each branch of the tree. The red triangles show the Costa Rican isolates. The forms of asexual sporulation of the aetiological agents of chromoblastomycosis, found in this study are: (A) branched conidiophore, (B, D) simple acroteca-type conidiophore and (C) phyalid-type conidiophore.

theses). The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) are shown next to the branches.

*F. pedrosoi* is the most frequently isolated species from chromoblastomycosis lesions worldwide [10,11,17,31,32], so it is not surprising that 60% of the isolates from this study were identified as this species. On the other hand, Yaguchi, *et al.* (2007) [17], reclassified a clinical isolate from Costa Rica from *F. pedrosoi* to *F. monophora* (GenBank access code AB117978 by ITS and AB253501 by cytochrome b sequencing), evidencing its presence in the country; data which was corroborated in this study. In turn, being Costa Rica a tropical country that has a high microbial diversity, it is expected that the other genera that cause this disease are also present. For example, Mora and collaborators reported in 2010 the first Costa Rican clinical case of chromoblastomycosis caused by *C. carrionii* [26] and we are currently describing the first case caused by *R. aquaspersa*.

Regarding the phylogenetic analysis, the tree obtained showed a separation of genera and species in clades with good statistical support (100% bootstrap). The retention index shows that the differences in the tree are in the internal nodes and that the isolates under study were generated from the same common ancestor.

When comparing the sequences of *Fonsecaea* spp. with those existing in the GenBank database, we found similarities with sequences from Latin American countries such as Brazil and Mexico. This phenomenon could be attributed to the possible similarity of ecosystems in tropical countries where the fungus can develop. However, in the case of *R. aquaspersa*, there is a greater relationship with sequences from Japan, Mexico and India. Finally, the molecular biology of pathogenic fungi has provided increasingly valuable information regarding the distribution and presence of previously unknown species.

### Conclusion

*F. pedrosoi*, *F. monophora* and *R. aquaspersa* were identified as etiological agents of chromoblastomycosis in Costa Rica. Hence, the standardization of a methodology to differentiate *Fonsecaea* species would represent a better clinical approach, since it would be known if the causative agent of the patients' disease has the capacity to disseminate or not.

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### Conflict of Interest

The authors declare that there is no conflict of interest involved in this article.

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