Introduction

Black grain eumycetoma represents the most common fungal mycetoma worldwide. This chronic, erosive infection of subcutaneous tissues particularly affects the lower extremities and leads to severe disability [1]. The disease is considered a major health problem in tropical areas and is prevalent among people of low socio-economic status [2].

Mycetoma presents as a subcutaneous mass with multiple sinuses that discharge pus, serous fluid and grains, i.e. the characteristic compact grains of the causative agent formed inside the lesion [3].

A wide range of microorganisms has been reported to cause mycetoma. For treatment, not only differentiation between (fungal) eumycetoma and (bacterial) actinomycetoma is important, but also the identity of the causative agent, since species differ in their response to antimicrobial drugs [4]. In endemic countries, clinical diagnosis may be the only diagnostic method. A fully developed mycetoma lesion is easily identified clinically, whereas in early stages with the absence of grains, the infection may be confused with phaeomycosis or soft tissue tumors [1]. In such cases fine needle aspiration cytology or deep surgical biopsy for histological examination are useful [1,5]. Some fungal and bacterial grains have a characteristic histological appearance which helps in provisional identification, but recognition of the causative species remains impossible [6]. Isolation of the pathogen from discharged grains or from biopsies allows identification of agents that sporulate, but most of the species lack phenotypic characteristics [3]. Molecular techniques have been introduced to facilitate the identification of nondescript organisms [7,8,9], but are of high cost and time-consuming. Thus, there is a need for a fast, simple and reliable method for identification.

Rolling circle amplification (RCA) is a powerful diagnostic method based on detection of specific nucleic-acid sequences and enzymatic amplification of circularized oligonucleotide probes under isothermal conditions [10]. The probes are linear oligonucleotides that contain two target-complementary sequences at their ends joined by linkers [11]. The ends of the probe hybridize to the complimentary target in juxtaposition and then ligate which results in series of repeats of the original circular template [10]. In addition, with the isothermal RCA assay, 62 isolates were successfully identified with 100% specificity and no cross reactivity or false results. The main advantage of this technique is the low-cost, high specificity, and simplicity. In addition, it is highly reproducible and can be performed within a single day.
mycetoma species *Scedosporium boydii* [18]. The aim of the present study is to develop RCA-based diagnostics for the most common agents of black-grain eumycetoma.

**Materials and Methods**

**Strains analyzed**

The study included 62 isolates belonging to eight species causing black grain mycetoma: *Madurella mycetomatis* (n = 32), *M. fahalii* (n = 1), *M. pseudomyxomycetotis* (n = 3), *M. tropicana* (n = 2), *Trematosphaeria grisea* (n = 10), *Faciosporum senegalensis* (n = 6), *F. tomkinsii* (n = 2), and *Medicopsis romeroi* (n = 6). Strains were obtained from the reference collections of CBS-KNAW Fungal Biodiversity Centre (Utrecht, The Netherlands) and the Mycetoma Research Centre (MRC, Khartoum, Sudan) and are listed with metadata in Table 1. Type strains of all tested species (*M. mycetomatis* (n = 1), *M. tropicana* (CBS 129176) DNA, 0.1 mM each dNTP, 0.6 U Taq polymerase (GC Biotech, 25 mM MgCl₂, 0.1% gelatin, 1% Triton X-100). PCR reactions were detected by electrophoresis using 1% agarose gels. The specificity of the 8 RCA probes was tested using strains of different fungal orders [19,20].

**DNA extraction and target amplification**

DNA was extracted using cetyltrimethylammonium bromide (CTAB) method as described by Möller et al. [21]. Amplification of the ITS region was performed using primers V9G and LS266 [22] in a 25 μL reaction mixture containing: 10 ng of template DNA, 0.1 mM each dNTP, 0.6 U Taq polymerase (GG Biotech, Alphen aan den Rijn, The Netherlands), 1 μL of each primer (10 pmol) and 2.5 μL reaction buffer (0.1 mM Tris-HCl, 0.5 M KCl, 25 mM MgCl₂, 0.1% gelatin, 1% Triton X-100). PCR reactions consisted of a 5 min pre-denaturation step at 95°C, followed by 30 cycles of 95°C for 30 s, 52°C for 30 s and 72°C for 1 min, with final post elongation step at 72°C for 7 min. PCR products were detected by electrophoresis using 1% agarose gels.

**Padlock probe design**

Sequences of the ITS region were used to design 8 probes specific for each species used in this study. Two alignments were generated since the analyzed species were known to belong to two different fungal orders [19,20]. ITS derived from *Madurella (Sordariales)* were aligned with 200 isolates of *Chaetomiaceae* including *Chaetomium, Thielavia,* and *Achatomum*. For the remaining species (*Pleosporales*) an alignment was constructed to include representative isolates of the family *Trematosphaeriaceae* and of coelomycetes in the suborder *Pleosporineae*. Sequences were aligned using BioNumerics v4.61 (Applied Maths, Sint-Martens-Latem, Belgium). Probes were designed with minimum secondary structure and were checked using PrimerSelect (DNASTAR Lasergene, WI, U.S.A.). To insure specificity of the probes, target-specific sites of each padlock probe was submitted to BLAST in NCBI sequence database for homologous sequences.

**Exonucleolysis**

Prior to RCA amplification reaction and in order to reduce the ligation-independent amplification, ligation products were treated by addition of 10 U exonucleases I and 10 U exonucleases III (New England Biolabs, Hitchin, U.K.) with a total reaction volume of 10 μL. Ligation conditions were: 5 min denaturation at 94°C, followed by 7 cycles of 94°C for 30 sec, 63°C for 4 min, and final cooling at 10°C.

**Rolling circle amplification (RCA)**

RCA amplification reaction was performed in a 50 μL mixture containing: 2 μL ligation product, 0 U E. coli DNA polymerase (New England Biolabs), 10 pmol of each RCA primer (Table 2), and 400 μM dNTP mix. The mixture was incubated at 65°C for 60 min and cooled at 10°C. Electrophoresis on a 1% agarose gel was used to visualize RCA products. A positive reaction is indicated by the presence of ladder-like pattern. The result was also visualized by adding 1.0 μL of a 10-fold diluted SYBR Green I (Cambrex BioScience, Workingham, U.K.) to 10 μL of the amplification product. Accumulated double stranded DNA was detected with UV transilluminator (Vilber Lourmat, Marne-la-Vallée, France).

**Determination of analytical specificity and sensitivity**

The specificity of the 8 RCA probes was tested using strains of black-grain mycetoma causative species listed in table 1. Analytical sensitivity was determined using 10-fold serial dilution of *M. mycetomatis* (CBS 109801) and *M. fahalii* (CBS 129176) DNA and the test was performed as mentioned above. In addition, RCA was performed directly using DNA samples without amplification of the target gene. To evaluate the detection limit from direct DNA samples two-fold serial dilutions of target DNA were tested. The sensitivity of the RCA probes was also determined by 10-fold serial dilution of MYC and MFAH probes tested with amplified ITS of *M. mycetomatis* and *M. fahalii* respectively.

**Results**

RCA was used to identify 62 strains belonging to eight species causing human eumycetoma. Since black grain eumycetoma species are known to be phylogenetically distant, it is easy to find unique sites for their identification. The ribosomal ITS region was sufficient for identification of all species and showed no intraspecific variability within a set of 100 *M. mycetomatis* strains in our collection. For *M. mycetomatis, M. tropicana, M. pseudomyxomycetotis,* and *F. senegalensis* the ITS1 region was...
| No. | Source | Origin          |
|-----|--------|-----------------|
| 1.  | Falciformispora senegalensis CBS 196.79 | Mycetoma Senegal |
| 2.  | Falciformispora senegalensis CBS 197.79 | Human Senegal    |
| 3.  | Falciformispora senegalensis CBS 198.79 | Mycetoma Senegal |
| 4.  | Falciformispora senegalensis CBS 199.79 | Human Senegal    |
| 5.  | Falciformispora senegalensis CBS 132257 | Mycetoma Sudan    |
| 6.  | Falciformispora senegalensis CBS 132272 | Mycetoma Sudan    |
| 7.  | Falciformispora tompkinsii CBS 200.79 | Mycetoma Senegal  |
| 8.  | Falciformispora tompkinsii CBS 201.79 | Mycetoma Senegal  |
| 9.  | Medicopsis romeroi CBS 252.60 | Mycetoma Venezuela |
| 10. | Medicopsis romeroi CBS 132878 | Mycetoma India |
| 11. | Medicopsis romeroi CBS 122784 | Plant |
| 12. | Medicopsis romeroi CBS 123975 | Phaeohyphomycosis India |
| 13. | Medicopsis romeroi CBS 128765 | Subcutaneous cyst Kuwait |
| 14. | Medicopsis romeroi CBS 135987 | onychomycosis Netherlands |
| 15. | Trematosphaeria grisea CWZ 29591 | |
| 16. | Trematosphaeria grisea CBS 332.50 | Mycetoma Chili |
| 17. | Trematosphaeria grisea CBS 246.66 | Submandibular abscess India |
| 18. | Trematosphaeria grisea CBS 120271 | Tap water The Netherlands |
| 19. | Trematosphaeria grisea CBS 135982 | Pastry gel The Netherlands |
| 20. | Trematosphaeria grisea CBS 135984 | Water The Netherlands |
| 21. | Trematosphaeria grisea CBS 136543 | Water The Netherlands |
| 22. | Trematosphaeria grisea CBS 135985 | Water The Netherlands |
| 23. | Trematosphaeria grisea CBS 135986 | Water The Netherlands |
| 24. | Madurella mycetomatis CBS 132258 (Mm10) | Mycetoma Sudan |
| 25. | Madurella mycetomatis CBS 132259 (Mm13) | Mycetoma Sudan |
| 26. | Madurella mycetomatis CBS 132260 (Mm14) | Mycetoma Sudan |
| 27. | Madurella mycetomatis CBS 132261 (Mm16) | Mycetoma Sudan |
| 28. | Madurella mycetomatis CBS 132262 (Mm18) | Mycetoma Sudan |
| 29. | Madurella mycetomatis CBS 132263 (Mm22) | Mycetoma Sudan |
| 30. | Madurella mycetomatis CBS 132265 (Mm28) | Mycetoma Sudan |
| 31. | Madurella mycetomatis CBS 132266 (Mm29) | Mycetoma Sudan |
| 32. | Madurella mycetomatis CBS 132267 (Mm30) | Mycetoma Sudan |
| 33. | Madurella mycetomatis CBS 132269 (Mm33) | Mycetoma Sudan |
| 34. | Madurella mycetomatis CBS 132270 (Mm36) | Mycetoma Sudan |
| 35. | Madurella mycetomatis CBS 132273 (Mm44) | Mycetoma Sudan |
| 36. | Madurella mycetomatis CBS 132274 (Mm45) | Mycetoma Sudan |
| 37. | Madurella mycetomatis CBS 132285 (Mm46) | Mycetoma Sudan |
| 38. | Madurella mycetomatis CBS 132276 (Mm49) | Mycetoma Sudan |
| 39. | Madurella mycetomatis CBS 132277 (Mm51) | Mycetoma Sudan |
| 40. | Madurella mycetomatis CBS 132284 (Mm54) | Mycetoma Sudan |
| 41. | Madurella mycetomatis CBS 131320 (Mm55) | Mycetoma Sudan |
| 42. | Madurella mycetomatis CBS 132280 (Mm58) | Mycetoma Sudan |
| 43. | Madurella mycetomatis CBS 132281 (Mm63) | Mycetoma Sudan |
| 44. | Madurella mycetomatis CBS 132282 (Mm64) | Mycetoma Sudan |
| 45. | Madurella mycetomatis CBS 132283 (Mm68) | Mycetoma Sudan |
| 46. | Madurella mycetomatis CBS 132284 (Mm71) | Mycetoma Sudan |
| 47. | Madurella mycetomatis CBS 132285 (Mm72) | Mycetoma Sudan |
| 48. | Madurella mycetomatis CBS 132286 (Mm73) | Mycetoma Sudan |
| 49. | Madurella mycetomatis CBS 132287 (Mm78) | Mycetoma Sudan |
| 50. | Madurella mycetomatis CBS 132288 (Mm83) | Mycetoma Sudan |
selected for probe design, while for \textit{M. fahalii}, \textit{T. grisea}, \textit{F. tompkinsii} and \textit{M. romeroi} the ITS 2 region was found to be more suitable.

RCA results for the tested strains were easily visualized in 1% agarose gel. Positive reactions demonstrated ladder like patterns while negative reactions resulted in a clear background (Fig. 1). With SYBR green, positive results showed green fluorescence when exposed to UV light, while negatives did not. When exonucleolysis was performed some inhibition was observed with low RCA positive signals on gel or with fluorescence. Faint non-specific bands were observed when this step was omitted. RCA reactions were performed successfully without digestion with exonucleases, as the non-specific bands did not interfere with RCA results. All \textit{M. mycetomatis} strains were correctly identified with RCA, irrespective of their geographical origin (Sudan, India, Mali) (Fig. 2). For the other agents, each individual species-specific probes yielded positive results with their corresponding species and with 100% agreement with ITS sequencing (Fig. 2, Table 3). No cross reactivity or false positive and negative results were observed.

The sensitivity of RCA when using amplified product of the target gene was less than $32 \times 10^{-3}$ ng of DNA. A higher concentration of 100 ng is needed when the test is carried out directly from the DNA samples without amplification of the ITS. The probes were very sensitive and a concentration of $6.6 \times 10^{-5}$ ng was successfully ligated and then amplified with RCA.

The turnaround time required for conducting the entire experiment including PCR amplification of target DNA, RCA

| Table 1. Cont. |

| Name | No. | Source | Origin |
|------|-----|--------|--------|
| \textit{Madurella mycetomatis} | CBS 109801 | Mycetoma | Sudan |
| \textit{Madurella mycetomatis} | CBS 110087 | Mycetoma | Sudan |
| \textit{Madurella mycetomatis} | CBS 110359 | Mycetoma | Mali |
| \textit{Madurella mycetomatis} | CBS 110356 | Mycetoma | Mali |
| \textit{Madurella mycetomatis} | CBS 132419 | Mycetoma | India |
| \textit{Madurella mycetomatis} | CBS 132589 | Mycetoma | India |
| \textit{Madurella tropicana} | CBS 201.38 | Mycetoma | Indonesia |
| \textit{Madurella tropicana} | CBS 331.50 | Mycetoma | New Mexico |
| \textit{Madurella pseudomyctomatis} | CBS 129177 | Mycetoma | China |
| \textit{Madurella pseudomyctomatis} | CBS 216.29 | Mycetoma | New Mexico |
| \textit{Madurella pseudomyctomatis} | CBS 248.48 | Mycetoma | New Mexico |
| \textit{Madurella fahalii} | CBS129176 | Mycetoma | Sudan |

(CBS Centraalbureau voor Schimmelcultures; Between brakets Erasmus collection number for strains from Sudan; Type strains marked with T)

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| Table 2. Oligonucleotide padlock probes and probe-specific primers used for species identification with RCA. |

| Species name | Probe and primer name | Sequences |
|--------------|-----------------------|-----------|
| \textit{M. tropicana} | MTROP | 5’pGAGACAAACAGGGTGTTGATAgatcaTGCTTCTCGTGCCCAATAcgcgggtcgcgtgtagtacCGCGCAGACACGATAgcttaAGAAGGGCCTAC3’ |
| \textit{M. pseudomyctomatis} | MPSREU | 5’pGAGACAAACAGGGTGTTGATAgatcaTGCTTCTCGTGCCCAATAcgcgggtcgcgtgtagtacCGCGCAGACACGATAgcttaAGAAGGGCCTAC3’ |
| \textit{M. fahalii} | MFAH | 5’pTGATACACTACGCTCGAGTGAAGGatcaTGCTTCTCGTGCCCAATAcgcgggtcgcgtgtagtacCGCGCAGACACGATAgcttaAGAAGGGCCTAC3’ |
| \textit{T. grisea} | TGRIS | 5’pACCCCTAGAGTCTCCCAAAAAGGCGatcaTGCTTCTCGTGCCCAATAcgcgggtcgcgtgtagtacCGCGCAGACACGATAgcttaAGAAGGGCCTAC3’ |
| \textit{F. senegalensis} | FSEN | 5’pATACAAGACAGGTTTGCCCGGatcaTGCTTCTCGTGCCCAATAcgcgggtcgcgtgtagtacCGCGCAGACACGATAgcttaAGAAGGGCCTAC3’ |
| \textit{F. tompkinsii} | FTOM | 5’pCTCTCACAAGTGCCCAAAAGGCGatcaTGCTTCTCGTGCCCAATAcgcgggtcgcgtgtagtacCGCGCAGACACGATAgcttaAGAAGGGCCTAC3’ |
| \textit{M. romeroi} | MRO | 5’pAGGCGAGGCTCCAGACACTCGTAGGatcaTGCTTCTCGTGCCCAATAcgcgggtcgcgtgtagtacCGCGCAGACACGATAgcttaAGAAGGGCCTAC3’ |
| \textit{M. mycetomatis} | MYC | 5’pACTACACTACGGGGAGGGGCGatcaTGCTTCTCGTGCCCAATAcgcgggtcgcgtgtagtacCGCGCAGACACGATAgcttaAGAAGGGCCTAC3’ |
| RCA1 | 5’-ATGGGCAACAGAGCA3’ |
| RCA2 | 5’-CCGGCAGACACGCTA3’ |

5’p- indicate phosphorylation of 5’ end, probes binding arms are underlined, the arms joined with non specific region lower case and RCA1 and RCA2 primer binding regions are bolded.

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RCA for Identification of Mycetoma Agents

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processing and analysis was found to be 6 hours. DNA sequencing of the ITS region took more than 8 hours to be performed (Fig. 3).

Discussion

Mycetoma is a unique tropical disease, endemic in many tropical and subtropical regions that has been recently added to the WHO list of neglected tropical diseases [24]. It is mainly prevalent in what is known as “mycetoma belt” which includes Mexico, Senegal, Sudan, India and other countries between tropic of cancer [1]. In 2014, a mycetoma consortium of scientists and physicians published research gaps on mycetoma which need to be addressed in the coming years [2]. One of the research priorities identified was the need to develop a reliable and cost-effective method for species identification to improve diagnosis [2].

Mycetoma agents have been extensively studied in recent years [8,9,20]. The large phylogenetic distance between a number of these agents provides the possibility to use a moderately variable marker like rDNA ITS for species identity. Ahmed et al. [25] developed PCR-restriction fragment length polymorphism (RFLP) for identification of M. mycetomatis targeting the ITS region. However, with the description of the molecular siblings M. fahalii, M. pseudomycetomatis, and M. tropicana [26] the method might be insufficiently accurate. Moreover, there is a need for identification these siblings species; Madurella grisea appeared to be distantly related and was re-named as T. grisea [20].

In the present study we developed a simple, fast and highly specific molecular method for the identification of agents of black grain mycetoma. In this method, the ITS region is easily amplified using one set of primers, which simplifies the use. In a second, isothermal amplification reaction padlock probes are used to identify the species by RCA. The only equipment necessary is a thermocycler for the PCR reaction and a water bath or heating block for the RCA reaction. This relative simplicity enhances possible use in routine laboratories in endemic areas. Due to its

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**Figure 1. Specificity of rolling circle amplification probes.** Agarose gel electrophoresis analysis of rolling circle amplification products. Positives probe signal seen as band pattern was only present with matched template–probe mixtures. Probe names are indicated on the top of the gel. Lanes; 1 M. mycetomatis CBS 109801, 2 M. tropicana CBS 201.38, 3 M. pseudomycetomatis CBS 129177, 4 M. fahalii CBS 129176, 5 T. grisea CBS 332.50, 6 F. senegalensis CBS 196.79, 7 F. tompkinsii CBS 200.70, 8 M. romeroi CBS 252.60, M DNA ladder.

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**Figure 2. Madurella mycetomatis identification by RCA.** Gel representation of rolling circle amplification reaction using Madurella mycetomatis probe (MYC) for strains recovered from mycetoma patient of origin: lane 1–18 Sudan; lane 19, 20 Mali; lane 21, 22 India; lane 23 negative control water; lane M ladder.

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robustness, high potential, and reproducibility, RCA is increasingly used as a diagnostic tool in pathogenic fungi, e.g., agents of chromoblastomycosis, dermatophytes, *Aspergillus*, *Candida*, and *Talaromyces marneffei* [16,23,27,28]. The method does not require DNA sequencing and is therefore considered as a rapid and cost-effective. Applications are being expanded to nano- and biotechnology [29].

In the present study eight species-specific probes were designed and used for identification of 62 isolates. For the RCA reaction species probe hybridization to the 3′ and 5′ ends of target DNA and joining of adjacent ends by DNA ligase when both show perfect complementarity. The ligation appears to be highly specific and thus the method can detect single nucleotide polymorphism [30]. The amplification reaction is driven by an isothermal DNA polymerase to amplify the circularized probes with high efficiency and an estimated capacity to synthesize more than 70,000 bp per hour [31]. RCA products can be detected with different methods including gel electrophoresis, radiolabeling, UV absorbance, fluorescence, and single molecule detection [32]. It was known that the positive signals can be detected within 15 min after starting the RCA reaction by real time PCR [23]. In the present study the RCA positive signal was easily visualized using both gel electrophoresis and fluorescent dye. The duration of our RCA protocol was 2 h, but additional time is required for DNA extraction and ITS amplification. Compared to the DNA sequencing the turnaround time for RCA is 2 hours less than sequencing and this even more if there is no in-house sequencer available.

Our results with eight padlock probes showed that RCA accurately identified all species with no cross reactivity (Fig. 1). It may be concluded that RCA is extremely useful for specific identification of agents of mycetoma. Performance and rapid turnaround time features make the RCA suitable for quick and reliable diagnosis, which is an enormous improvement compared to the current phenotypic identification of mostly non-sporulating cultures. Future application of RCA could be the detection of agents DNA directly from clinical samples without requirement of culturing.

### Supporting Information

**S1 Figure** STARD flowchart for RCA. (PDF)

**Table 3. Rolling circle amplification results of analysed strain.**

| Strains   | M. mycetomatis (3) | M. tropicana (2) | M. pseudomycetomatis (3) | M. fahalii (1) | T. grisea (10) | F. senegalensis (6) | F. tompkinsii (2) | M. romeroi (6) |
|-----------|--------------------|------------------|--------------------------|----------------|----------------|----------------------|------------------|---------------|
| Positive results (+), negative results (−). | + | − | − | − | − | − | − | + |

Positive results (+), negative results (−).

Figure 3. Identification time of species using rolling circle amplification (RCA) and sequencing of ITS.
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Conceived and designed the experiments: SAA BHGGvdE GSdH. Performed the experiments: SAA. Analyzed the data: SAA BHGGvdE. Contributed reagents/materials/analysis tools: GSdH AHF WWJvdS. Wrote the paper: SAA BHGGvdE WWJvdS GSdH AHF.
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