Development and Validation of a Multiplex PCR for Detection of \textit{Scedosporium} spp. in Respiratory Tract Specimens from Patients with Cystic Fibrosis

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The emergence of \textit{Scedosporium} infections in diverse groups of individuals, which are often treatment refractory, warrants timely and accurate laboratory diagnosis. Species- or group-specific primers based on internal transcribed spacer (ITS) sequence polymorphisms were designed for \textit{Scedosporium aurantiacum}, \textit{Scedosporium dehoogii}, \textit{Scedosporium prolificans}, \textit{Pseudallescheria boydii} species complex (former clade 5)/\textit{Pseudallescheria apiosperma} (formerly classified as \textit{S. apiospermum} sensu lato) and \textit{Pseudallescheria minutispora}. Primers for \textit{S. aurantiacum}, \textit{S. prolificans}, and \textit{P. boydii} species complex/\textit{P. apiosperma} were incorporated into a multiplex PCR assay for the detection and identification of the three major clinically important \textit{Scedosporium} species and validated using sputum specimens collected from patients seen at a major Australian cystic fibrosis clinic. The multiplex PCR assay showed 100% specificity in identifying the three major clinically relevant \textit{Scedosporium} species from pure culture. When evaluated using DNA extracts from sputa, sensitivity and specificity of the multiplex PCR assay were 62.1% and 97.2%, respectively. This highly species-specific multiplex PCR assay offers a rapid and simple method of detection of the most clinically important \textit{Scedosporium} species in respiratory tract specimens.

\textit{Scedosporium} species are emerging fungal pathogens capable of causing a wide range of infections, particularly in seriously ill and immunocompromised patients (8, 26). Long-term colonization with these fungi has been reported in patients with structurally abnormal respiratory airways, including those with cystic fibrosis (CF) (1, 2, 6, 9). Invasive fungal infection is rare in CF patients prior to lung transplantation (15), yet \textit{Scedosporium} colonization may be a risk factor for invasive infection posttransplantation. Since \textit{Scedosporium} spp. are resistant to many antifungal agents, colonization is a relative contraindication for transplantation in some centers (8, 24).

Among clinical isolates, the most frequently encountered \textit{Scedosporium} species include \textit{Scedosporium prolificans}, \textit{Scedosporium apiospermum}/\textit{Pseudallescheria apiosperma}, \textit{Pseudallescheria boydii} sensu stricto, and the recently described species \textit{Scedosporium aurantiacum} (11, 13, 14). The emergence of \textit{Scedosporium} infections in Australia has been highlighted in earlier reports describing the species distribution, clinical epidemiology, and outcomes (1, 2, 7, 10, 16). Further, a nationwide population-based study recovered a substantial number of infections due to \textit{S. aurantiacum} and identified an association between isolation of this species and the presence of chronic lung disease (16). Invasive fungal infection is associated with significant morbidity and mortality rates of up to 75 to 80% (17, 25, 26). Therefore, early and accurate detection of \textit{Scedosporium} infections is crucial for prompt and effective treatment.

Conventional mycological methods for detection and identification of \textit{Scedosporium} spp. in clinical specimens, however, are insensitive and time-consuming, as culture from sputum samples may require up to 14 days to produce fungal growth adequate for morphological identification (1, 5). A number of molecular detection and identification methods for \textit{Pseudallescheria}/\textit{Scedosporium} have been reported (3, 4, 19, 20, 21, 29). Real-time PCR protocols for detection of \textit{S. prolificans} and \textit{S. apiospermum} sensu lato have been developed (4). Bouchara et al. employed an oligonucleotide array targeting the internal transcribed spacer (ITS) region of the ribosomal DNA (rDNA) gene cluster for direct detection of a broad range (\textit{n} = 20 species) of fungal pathogens, including the \textit{Pseudallescheria}/\textit{Scedosporium} species complex and \textit{S. prolificans} in the sputum of CF patients (3). An ITS-directed pan-fungal PCR assay combined with DNA sequencing was used to detect multiple fungal genera, including \textit{S. prolificans}, in fresh and formalin-fixed clinical specimens (19). Recently, multiplex-tandem PCR (MT-PCR)- and rolling circle amplification (RCA)-based approaches successfully identified \textit{S. prolificans} and \textit{S. apiospermum} sensu lato from isolated colonies (both methods) and from blood cultures (MT-PCR) (20, 21, 29). However, most of these assays do not take into account the taxonomic reclassi-
fication of *Scedosporium* spp. or are designed to detect only a few *Scedosporium* species. Further, they required either an additional sequencing step, which may pose a problem in the case of mixed infections, or need specialized equipment, which may prevent them from being easily implemented into the routine microbiology laboratory in the near future.

In the present study, species-specific primers for the most common *Scedosporium* species (*S. aurantiacum*, *S. prolificans*, and *P. boydii* species complex/*P. apiosperma*) were designed, and a multiplex PCR method was developed to enable simultaneous rapid and accurate detection and identification of the major pathogenic *Scedosporium* species directly from clinical samples. The performance of the assay was then evaluated using sputum specimens from adult patients with CF.

### MATERIALS AND METHODS

**Fungal isolates.** To develop and validate the herein-reported species-specific primers and multiplex PCR protocol, the following fungal cultures were used: *S. prolificans* (strains WM 06.399 and FMR 7248), *P. apiosperma* (strains WM 06.482 and FMR 8630), *S. dehoogii* (strain FMR 6921), *P. boydii* (strain FMR 4167), *S. minitanspora* (strain FMR 4072), *S. aurantiacum* (strains WM 08.29), *P. minutispora* (strain WM 08.194 (FMR 8619, CBS 117407), *S. aurantiacum* (strains WM 06.355), *P. apiosperma* (strain WM 08.202) (FMR 8632, CBS 116910), and *S. prolificans* (strain WM 09.399) was used as a positive control in each reaction batch. The PCR products were detected as described above. The lowest DNA concentration (10 ng/μl) of sputum extract was used as a negative control, and genomic DNA extracted from *P. apiosperma* strain WM 08.194 (FMR 8619, CBS 117407), *S. aurantiacum* strain WM 08.202 (FMR 8630, CBS 116910), and *S. prolificans* strain WM 09.399 was used as a positive control in each reaction batch. The PCR amplification was performed in a thermal cycler (Perkin Elmer Cetus, Norwalk, CT) using the following conditions: an initial denaturation at 94°C for 5 min, followed by 35 cycles of 94°C for 45 s, 65°C for 45 s, and 72°C for 2 min, with each extension step at 72°C for 10 min. The PCR products were separated on 1.5% agarose gels in Tris-borate-EDTA (TBE) buffer and visualized by UV transillumination.

**Assay detection limit.** DNA extract from cultured *S. aurantiacum* strain WM 08.202 was serially diluted in sterile water. The PCR assay was performed as described above using a DNA template concentration from 1 fg to 1 ng per PCR. The PCR products were detected as described above. The lowest DNA concentration that produced a visible band was considered the detection limit of this assay.

### RESULTS

**Single PCR.** Using extracted DNA from pure cultures as a template, the individual single-species-specific PCRs produced species-specific band sizes (Table 1, Fig. 1). No cross amplification of any of the species-specific primers with the other *Scedosporium* species was observed (data not shown). Figure 2

<table>
<thead>
<tr>
<th>Name</th>
<th>Primer sequence</th>
<th>Amplicon size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. boydii</em> species complex</td>
<td>5′ AGGCCGTGCGGCTAACACCTAAC 3′</td>
<td>300</td>
</tr>
<tr>
<td><em>P. apiosperma</em></td>
<td>5′ CTACGCGTCTGCGCTGA 3′</td>
<td>300</td>
</tr>
<tr>
<td><em>P. minitanspora</em></td>
<td>5′ CTCGTTTGCGTTAGCGAAAGCTCAG 3′</td>
<td>450</td>
</tr>
<tr>
<td><em>S. aurantiacum</em></td>
<td>5′ TACGCACTCGTCGAAGGAGC 3′</td>
<td>450</td>
</tr>
<tr>
<td><em>S. dehoogii</em></td>
<td>5′ CGCCCGAAAGGAGCAGG 3′</td>
<td>650</td>
</tr>
<tr>
<td><em>S. prolificans</em></td>
<td>5′ CTACGCGTCTGCGCTGA 3′</td>
<td>650</td>
</tr>
</tbody>
</table>

**Multiplex PCR.** The multiplex PCR was performed using the same protocol as was used for the individual PCRs described above, with an equimolar amount of each of the three primer sets (listed in Table 1) added to the reaction mixture. The PCR amplifications were done using the same cycling conditions as described above. The resulting amplicons were separated on 1.5% agarose gels and visualized by UV transillumination.

**DNA extraction.** Extraction of genomic DNA was performed as previously described (22).

**Sputum sampling and extraction.** Two hundred and eight expectorated sputum samples were obtained from 69 patients attending the Westmead Hospital Adult CF Clinic, Westmead, NSW, Australia, from April 2008 to March 2009. Approval for the study was obtained from the human ethics review board of the Sydney Medical School—Westmead Hospital, Sydney, Australia, and subcultured onto Sabouraud dextrose agar (Oxoid, Hampshire, United Kingdom) at room temperature for 5 days to ensure adequate growth and purity prior to use.

**RESULTS**

**Single PCR.** Using extracted DNA from pure cultures as a template, the individual single-species-specific PCRs produced species-specific band sizes (Table 1, Fig. 1). No cross amplification of any of the species-specific primers with the other *Scedosporium* species was observed (data not shown). Figure 2

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Candida parapsilosis strain WM 04.547; 11, Aspergillus fumigatus prolife-
Scedosporium prolificans strain WM 06.482; 7, S. prolificans amplified with the primer pair MSPF1/MSPR2; M, 1-kb-plus DNA marker (Invitrogen, Carlsbad, CA).

tus

All Scedosporium spp. (n = 29)

62.1 (63.6) 97.2 (98.3) 78.3 (87.5) 94.1 (93.4)

TABLE 2. Test performance parameters of the PCR based on the number of specimens (and number of patients)

<table>
<thead>
<tr>
<th>Species</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>Positive predictive value</th>
<th>Negative predictive value</th>
</tr>
</thead>
<tbody>
<tr>
<td>S. aurantiacum (n = 15)</td>
<td>60.0</td>
<td>100</td>
<td>100</td>
<td>97.0</td>
</tr>
<tr>
<td>S. prolificans (n = 10)</td>
<td>70.0</td>
<td>99.0</td>
<td>77.8</td>
<td>98.5</td>
</tr>
<tr>
<td>P. boydii/S. apiosperma (n = 4)</td>
<td>50.0</td>
<td>98.5</td>
<td>40.0</td>
<td>99.0</td>
</tr>
<tr>
<td>All Scedosporium spp. (n = 29)</td>
<td>62.1</td>
<td>97.2</td>
<td>78.3</td>
<td>94.1</td>
</tr>
</tbody>
</table>

The multiplex assay had an excellent specificity for the three targeted Scedosporium species, with accurate species identification in all cases (data not shown). Multiplex PCR resulted in positive results in five culture-negative samples. The specificity of the developed multiplex PCR assay was 100% when performed on pure culture and 97.2% when performed on sputum extracts (Table 2), with a positive predictive value (PPV) of 78.3% and a negative predictive value (NPV) of 94.1% (Table 2). When the test was analyzed according to the number of patients, no significant changes in the obtained values were observed (see values given in parentheses in Table 2).

Bacterial and non-Scedosporium fungal cocolonization had no significant influence on the proportion of false-positive or false-negative samples; bacterial colonization and non-Scedosporium fungal colonization were identified in 13/18 (72.2%) and 13/18 (72.2%) of true positives, 5/5 (100%) and 2/5 (40%) primer pairs (data not shown). The assay yielded 100% specificity when tested on DNA extracted from pure fungal cultures with a detection limit of 100 fg of DNA (data not shown).

**Multiplex PCR.** In the multiplex PCR, no overlapping amplifications between the different primer pairs of S. aurantiacum, S. prolificans, and P. boydii species complex/P. apiosperma were obtained (Fig. 2). The same species-specific bands were generated as had been obtained in the individual PCRs for each species (data not shown). However, cross-amplification occurred between the S. dehoogii- and P. minutispora-specific primer pairs with the other Scedosporium species-specific primers when used in a multiplex setting, leading to the exclusion of these species from the multiplex PCR assay.

The multiplex PCR assay was validated using DNA extracted from 208 respiratory samples (sputa) from 69 patients. All positive results were repeated at least twice to exclude human or technical error. The sensitivity and specificity of the assay were calculated against the identification results obtained from culture as the gold standard. Nonselective and/or selective culture isolated Scedosporium spp. in 29 sputum samples (13.9%; S. aurantiacum, 15 samples; S. prolificans, 10 samples; P. boydii species complex/P. apiosperma, 4 samples) from 11 patients. Scedosporium spp. were detected by PCR in 18 of the 29 culture-positive specimens (sensitivity, 62.1%) (Table 2) and in 7 of the 11 patients (sensitivity, 63.3%). Eleven false-negative PCRs were noted from four patients; these included six samples from one patient.

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Bacterial and non-Scedosporium fungal cocolonization had no significant influence on the proportion of false-positive or false-negative samples; bacterial colonization and non-Scedosporium fungal colonization were identified in 13/18 (72.2%) and 13/18 (72.2%) of true positives, 5/5 (100%) and 2/5 (40%)
of false positives, 11/11 (100%) and 11/11 (100%) of false negatives, and 30/174 (17.2%) and 141/174 (81.0%) of true negatives. Specimen processing time (after sputum storage for less than 6 months or more than 6 months) had no significant influence on the proportion of false-positive or false-negative samples (data not shown).

**DISCUSSION**

_Scedosporium_ species are increasingly important fungal pathogens. Timely and accurate detection of these infective agents in clinical specimens is critical to the appropriate management of affected patients. PCR-based detection assays for pathogenic fungi have been widely used. It is important for these assays to be robust and to be easily performed in the routine mycology laboratory. The present study describes the design of five species-specific PCR primers and the subsequent development and validation of a multiplex PCR assay using three of the five species-specific primer pairs to detect and identify clinically important _Pseudallescheria/Scedosporium_ species in respiratory (sputum) specimens. The ITS region was used as the basis for the primer design due to its presence as a species in respiratory (sputum) specimens. The ITS region was important, as it is adequately variable to enable species differentiation within the _Pseudallescheria/Scedosporium_ species complex. The multiplex PCR assay showed a sensitivity and specificity of 100% using DNA from pure cultures, with clear and accurate differentiation of the three major pathogens—_S. prolificans, P. boydii_ species complex/ _P. apiosperma_, and _S. aurantiacum_. Other common fungal pathogens were not amplified, thus demonstrating the high specificity of this assay.

To evaluate the performance and clinical applicability of this assay, we tested its ability to detect and identify _Scedosporium_ spp. in a large number (n = 208) of sputum specimens collected from CF patients. These patients were studied since they have a propensity to be colonized by filamentous fungi, including _Scedosporium_ species (6, 9, 28), thus providing a suitable test population for the assay evaluation. When tested on expectorated sputum sample extracts, the multiplex PCR assay correctly identified the targeted _Scedosporium_ species with an overall sensitivity of 63.61% in the studied patients. This result is comparable to those found for PCR assays developed for the direct detection of fungal pathogens in clinical samples. For example, an _Aspergillus_ PCR assay showed a sensitivity ranging from 40% to 100% when performed on clinical specimens, including blood, bronchoalveolar lavage (BAL) fluid, fine-needle aspiration, or biopsy specimen (18). In a pan-fungal PCR assay including the detection of _S. prolificans_, the sensitivities were reported to be either 97% or 68% when applied on fresh tissue specimen or paraffin-embedded specimen, respectively (19). However, a study using an oligonucleotide array assay for the direct detection of fungi in sputum samples from a similar group of patients showed a sensitivity and specificity of 100% and 99%, respectively (3), which is most likely due to the different assay format.

Reviewing the false-negative PCRs, we noted that six of these samples, which grew _S. aurantiacum_ on culture, were serial samples belonging to the same patient. The possible factors contributing to the persistent negative reactions in this particular patient are unknown. An unexplainable PCR negativity was also observed for culture-positive BAL specimens in a previously reported study (18). Possible explanations for these and the other five false-negative results could include insufficient amounts of _Scedosporium_ DNA in the extracted sputa or the presence of endogenous PCR inhibitors. Inhibition of PCR in sputum samples following certain DNA isolation procedures has been reported previously for other PCR-based assays (23, 27).

The overall specificity of this assay in the patient cohort studied was 98.3% (Table 2), whereas the positive predictive value (PPV) and the negative predictive value (NPV) were 87.5% and 93.4%, respectively. False-negative results for the detection of _S. apiosperma_ were noted in five clinical samples, of which four samples belonged to a patient who had positive cultures in other samples (not subjected to PCR amplification, due to an insufficient amount of sputum). Nevertheless, we noted earlier that the assay specificity was 100% when used on DNA extracted from pure cultured isolates. The high specificity provides a great advantage as to specifically identifying _S. prolificans_, _P. boydii_ species complex/ _P. apiosperma_, and _S. aurantiacum_. This is particularly important, as it will help in confirming the identities of phenotypically atypical strains, a phenomenon not uncommonly observed among the clinical isolates in this study. Species identification is further important, as species-specific clinical associations and antifungal susceptibilities have been described (12, 16). The ability to detect the three major clinically important _Scedosporium_ species directly in sputum samples will substantially reduce the turnaround time for microbiological species identification, which will in turn allow a more timely response and better patient management.

In conclusion, this study developed species-specific primer pairs to be used either individually or in a multiplex PCR assay for the identification of the major clinically relevant _Scedosporium_ species from pure culture and for the direct detection in sputum samples. The herein-developed PCR assay can be performed in routine clinical laboratories with a basic PCR setup. This single-step assay allowed for a high analytical sensitivity with a detection limit of as low as 100 fg _Scedosporium_ DNA and with a high specificity. The assay can be used as a complement to current culture and morphological-based identification methods to reduce the turnaround time and to increase the detection rate. Even though the assay was evaluated on CF samples, it could equally be applied to other patient groups. Further evaluation of this assay using other clinical specimens, including blood and tissue, is indicated to determine its position as a diagnostic tool.

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REFERENCES


