Characterization and Comparison of Galactomannan Enzyme Immunoassay and Quantitative Real-Time PCR Assay for Detection of *Aspergillus fumigatus* in Bronchoalveolar Lavage Fluid from Experimental Invasive Pulmonary Aspergillosis


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Received 28 December 2006/Returned for modification 25 January 2006/Accepted 1 May 2006

Bronchoalveolar lavage (BAL) is widely used for evaluation of patients with suspected invasive pulmonary aspergillosis (IPA). However, the diagnostic yield of BAL for detection of IPA by culture and direct examination is limited. Earlier diagnosis may be facilitated by assays that can detect *Aspergillus* galactomannan antigen or DNA in BAL fluid. We therefore characterized and compared the diagnostic yields of a galactomannan enzyme immunoassay (GM EIA), quantitative real-time PCR (qPCR), and quantitative cultures in experiments using BAL fluid from neutropenic rabbits with experimentally induced IPA defined as microscopically and histologically evident invasion. The qPCR assay targeted the rRNA gene complex of *Aspergillus fumigatus*. The GM EIA and qPCR assay were characterized by receiver operator curve analysis. With an optimal cutoff of 0.75, the GM EIA had a sensitivity and specificity of 100% in untreated controls. A decline in sensitivity (92%) was observed when antifungal therapy (AFT) was administered. The optimal cutoff for qPCR was a crossover of 36 cycles, with sensitivity and specificity of 80% and 100%, respectively. The sensitivity of qPCR also decreased with AFT to 50%. Quantitative culture of BAL had a sensitivity of 46% and a specificity of 100%. The sensitivity of quantitative culture decreased with AFT to 16%. The GM EIA and qPCR assay had greater sensitivity than culture detection of *A. fumigatus* in BAL fluid in experimentally induced IPA (P ≤ 0.04). Use of the GM EIA and qPCR assay in conjunction with culture-based diagnostic methods applied to BAL fluid could facilitate accurate diagnosis and more timely initiation of specific therapy.

Invasive pulmonary aspergillosis (IPA) is a major cause of morbidity and mortality in immunocompromised patients (7–9, 13, 14, 18, 27). Mortality rates of this severe infection range from 30% to 90% in neutropenic patients (9, 10, 15, 20). Accurate diagnosis of IPA routinely relies upon bronchoalveolar lavage (BAL) as a standard of care in assessment of immunocompromised patients (1, 12). However, current culture-based methods for evaluation of BAL fluid may have low sensitivity (3, 19, 32, 33). Other methods, such as tissue biopsy, are often not optimal due to the fragile condition of the patient with respect to sustaining an invasive procedure. Early diagnosis of IPA remains difficult. Improved prognosis for IPA requires early diagnosis (40). The development of nonculture and noninvasive methods may facilitate an early diagnosis of IPA (5, 22, 25).

The sandwich enzyme immunoassay (EIA) based on the detection of the *Aspergillus* antigen galactomannan (GM) (3, 4, 21, 38) and real-time PCR methods (25, 34) for the detection of *Aspergillus*-specific DNA are encouraging alternatives to biopsy and culture. However, the sensitivity and optimal interpretive cutoff values of the GM EIA in studies using BAL fluid are not well defined (4, 34, 38). Although quantitative real-time PCR (qPCR) assays have been developed for the detection of IPA (16, 26, 34, 35), their diagnostic sensitivity and specificity for IPA in BAL fluid are not well understood. There is also a need for methods that will complement each other for the early detection of aspergillosis. We therefore hypothesized that the GM EIA and qPCR assay would be more sensitive than culture methods in detection of *A. fumigatus* in BAL fluid. In order to test this hypothesis, we studied the application of a qPCR assay for the detection of *Aspergillus* DNA in conjunction with the GM EIA for the detection of GM antigen in BAL fluid from experimentally induced IPA in persistently neutropenic rabbits. Receiver operator curves (ROC) were constructed for both GM EIA and qPCR. These curves were then used to define cutoff, sensitivity, and specificity values that optimized the test performance of each assay. Both methods were studied using a well-established persistently neutropenic rabbit model of IPA (11, 28, 42). We further investigated the impact of different antifungal modalities, including ravuconazole, micafungin, and deoxycholate amphotericin B, on the sensitivity and specificity of each assay.

**MATERIALS AND METHODS**

Animal model of IPA. (i) Animals. Healthy female New Zealand White rabbits (Hazleton Research Products, Inc., Denver, PA) weighing 2.6 to 3.5 kg at the...
time of inoculation were used in all experiments. Rabbits were monitored under conditions of humane care and use in facilities accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International and according to National Institutes of Health guidelines for animal care and guidelines of the National Research Council (6). Vascular access was established in each rabbit by the surgical placement of a silastic tunneled central venous catheter as described in a previous publication (41).

(ii) Organism and inoculation. Invasive pulmonary aspergillosis was established as described elsewhere (11). Briefly, Aspergillus fumigatus (A. fumigatus isolate NIH 4215, ATCC MYA-1163) obtained from a fatal case of IPA was used in all experiments. The concentration was adjusted to give each rabbit a predetermined endotracheal inoculum of 1 x 10^6 to 1.25 x 10^6 conidia of A. fumigatus in a volume of 250 to 350 l. The concentration of the inoculum was confirmed by plating serial dilutions on 5% Sabouraud glucose agar plates (Biotwoks, Inc., Baltimore, MD). Inoculation was performed on day 2 of the experiments under conditions of general anesthesia. The A. fumigatus inoculum was administered endotracheally under direct visualization with a tuberculin syringe attached to a 16 gauge, 5/4 in. Teflon catheter (Becton Dickinson Infusion Therapy Systems Inc., Sandy, UT).

(iii) Immunosuppression and maintenance of neutropenia. Immunosuppression and neutropenia were established as previously described (28).

(iv) Antifungal compounds and treatment groups. In order to study the effect of antifungal therapy on the possible diminution of sensitivity of GM EIA, qPCR, and culture, we studied compounds from each major class: triazoles, echinocandins, and polyenes. Antifungal therapy with voriconazole (BMS-379224; Bristol-Myers Squibb Pharmaceutical Research Institute, Princeton, NJ) was administered at dosages of 5 mg/kg body weight. Micafungin (Funigene Pharmaceutical Co., Ltd, Osaka, Japan) was administered at dosages of 0.5 to 2 mg/kg. Decoycholate amphotericin B (Fungizone; Bristol-Myers, Squibb Company, Princeton, NJ) was administered at 1 mg/kg. Antifungal therapy was administered for 12 days starting 24 h after endotracheal inoculation of A. fumigatus conidia.

Rabbits were euthanized according to Animal Care and Use Committee-approved prespecified humane end points by intravenous administration of pentobarbital (65 mg of pentobarbital sodium/kg; pentobarbital sodium was in the form of 0.5 ml of euthanasia solution) (Beuthanasia-D special; Schering-Plough Animal Health Corp., Union, NJ) at the end of each experiment. BAL was then performed at the time of postmortem examination.

(v) Tissue quantification of pulmonary aspergillosis. Rabbits were considered to have invasive pulmonary aspergillosis when pulmonary lesions, histology, and quantitative culture of lung tissue were detected as described previously (28).

Diagnostic assays. (i) Bronchoalveolar lavage and BAL fluid fungal cultures. A BAL was performed on each postmortem resected lung preparation by the instillation and subsequent withdrawal of 10 ml of sterile normal saline solution into the clamped trachea with a sterile 12 ml syringe. This process was repeated for a total infusion of 20 ml of normal saline. The lavage material was then centrifuged for 10 min at 200 x g. The supernatant was discarded, leaving 2 ml of BAL fluid in which the pellet was then resuspended. A 100 µl volume of this fluid and 100 µl of a dilution (10^-1) were cultured on Sabouraud glucose agar plates and incubated at 37°C for the first 24 h and then at room temperature for another 24 h. CFU per ml were counted, and log CFU/ml was calculated. These samples were obtained from animals used for other primary studies (29, 30). BAL stores were stored at -70°C and later studied for expression of GM antigen and detection of nucleic acid.

(ii) EIA for detection of galactomannan in BAL fluid and serum. GM antigen levels were determined by the Platelia Aspergillus EIA one-stage immunoenzyme-magnetic sandwich microplate assay method (Bio-Rad Laboratories, Redmond, WA). The assay was performed according to the manufacturer’s directions. The assay uses the rat monoclonal antibody EBA-2, which is directed against the Aspergillus GM molecule (36). The optical absorbance of specimens and controls was determined by use of a microplate spectrophotometer equipped with 450 nm and 620 nm filters (Multiscan MMC/340; Titertek, Huntsville, AL).

Quantitative real-time PCR assay. (i) DNA isolation from BAL fluid. To avoid potential contamination, DNA samples were extracted in an AirClean PCR workstation (AirClean Systems, Raleigh, NC). Frozen BAL fluid samples were brought to room temperature prior to DNA extraction. Samples were vortexed for 1 min, and 500 µl was centrifuged for 10 min at 16,000 x g; supernatants were then discarded. The pellets were gently resuspended in 100 µl xerophlan buffer (1.0 M potassium acetate [Sigma S-86018], 50.0 mM sodium phosphate monobasic [Sigma S-7051], 0.1% 2-mercaptoethanol ([Sigma M-3148], 10 mg/ml lyticase [Sigma L-2524]) and 10 µl of 10x lysing enzymes (Novozyme [Sigma L-1412]) (20 mg/ml) and incubated at 37°C on a rocking platform for 1 h. After centrifugation for 20 min at 400 x g, the xerophlan-BAL fluid pellets were resuspended in 400 µl API buffer (DNeasy Plant Mini kit; Qiagen). The samples were added to Lysing Matrix D tubes (BioPulverizer system I; Qiogene/MP Biomedical, Morgan Irvine, CA) and processed using a FastPrep instrument (Qiogene/MP Biomedical, Morgan Irvine, CA) (24). Samples were processed at speed 5 for 30 s and placed on ice for 5 min; this process was performed a total of three times. Samples were centrifuged at 16,000 x g for 60 s and then gently vortexed. The supernatants (approximately 300 µl) were transferred to new tubes. The beads in the Lysing Matrix D tubes were washed with 300 µl API buffer, and this wash was added to the samples (resulting in a 400 µl final volume). Four microliters of RNase A (100 mg/ml) was added, and the mixture was vortexed vigorously and incubated for 10 min at 65°C in an Eppendorf thermomixer (Eppendorf, Westbury, NY) at 1,200 rpm. The pellets were processed according to the protocol for the DNeasy Plant Mini kit (Qiagen, Valencia, CA) as described in a previous publication (26).

(ii) Threshold for detection of extrinsic contamination with Aspergillus conidia. A 500 µl aliquot of normal BAL fluid was spiked with 10^3, 10^4, or 10^5 conidia or was left unspiked, and DNA was extracted as described above. In addition, water was processed through the extraction protocol as a kit blank control. All samples were processed in triplicate. The A. fumigatus-specific qPCR assay used in these studies was able to detect ≥100 CFU of conidia in 500 µl of a normal BAL sample. Thus, the kit blank control would be able to detect any extrinsic contamination from environmental conidia at a threshold of ≥100 CFU. In order to design the primer and probe set, the qPCR assay was described in a previous publication (26). Briefly, a quantitative real-time PCR assay targeting the internal transcribed spacer 1, 5.8S, and internal transcribed spacer 2 regions of the rRNA gene complex and fluorescence resonance energy transfer technology were used. The NCBI BLAST database search program documented the specificity of the primers and probes for A. fumigatus (12).

(iv) qPCR conditions. The PCR master mix consisted of 0.5 µM concentration of each of the primers Cap (+) sense (5’ CCGAAGCCCGGAATTG 3’), and Cap (-) sense 5’TGGAGGCGGATTGC 3’, 5 µM MgCl2, 0.025% bovine serum albumin (Sigma-Aldrich Corp., St. Louis, MO), 0.025 1U/ml Platinum Taq DNA polymerase (Invitrogen Corp., Carlsbad, CA), PCR 10x buffer (Invitrogen Corp., Carlsbad, CA), 0.2 mM PCR Nucleotide Mix PLUS (Roche Molecular Biochemicals, Indianapolis, IN) (1 dATP, dCTP, dGTP, and 3 dUTP in proportionate ratios), and 0.1 µM each of the fluorescens (5’ AGTATGCGATCCTAGAGGTTATGC 3’ and LC Red-640 5’ ATCAGTAAAATTCTC AACACGGA 3’) probes. To prevent potential amplicon carryover, each reaction also contained HK-UNG thermostable uracil glycosylase (Epicenter, Madison, WI) as recommended by the manufacturer. Each reaction contained a 5 µl aliquot of extracted specimen together with 15 µl of the master mix. A LightCycler 2.0 instrument (Roche Applied Science, Indianapolis, IN) was used with the following cycling conditions: uracil activation, 37°C for 180 s; uracil heat inactivation, 95°C for 10 s for 1 cycle. Amplification cycles were as follows: denaturation at 95°C for 10 s (20°C), annealing at 55°C for 10 s (20°C), annealing at 72°C for 1 s (20°C), extension at 72°C for 1 s (3°C), and cool down at 40°C for 120 s. The total number of cycles was 45. Quantitation standards (serial dilutions of A. fumigatus genomic DNA ranging from 5 x 10^5 fg to 5 x 10^1 fg) were run in conjunction with each set of samples. The following controls were also included: DNA extracted from normal BAL fluid, kit blank control (water processed through extraction protocol), and negative master mix control (water). All samples were run in triplicate, and the median cycle number data were used for analysis. The master mix was prepared in a biosafety cabinet in a different room from that in which DNA extractions were performed. LightCycler carousel loading was performed in a separate room from that in which the PCR master mix was prepared.

(v) Analytical sensitivity and quantitation of the qPCR assay. Quantitation, accuracy, and precision of the qPCR assay were determined through serial dilutions of A. fumigatus genomic DNA ranging from 5 x 10^5 fg to 5 x 10^1 fg. The calculated PCR efficiency was 1.98. A quantitative PCR signal of <3 cycles was considered a positive result. Fluorescent curves were analyzed with LightCycler software, version 3.5. The analysis of quantification data was done by the second derivative maximum method. This highly reproducible method performs automatic data calculations and determines crossing points with no user influence.

(vi) Specificity. The specificity of the assay was determined by cross-reactivity studies using 100 fg of genomic DNA from the following organisms: Aspergillus flavus, Aspergillus niger, Aspergillus terreus, Penicillium marneffei, Penicillium notatum, Penicillium citrinum, Penicillium purpureum, Pseudallescheria boydii, Penicillium chrysogenum, Rhizopus oryzae, Fusarium solani, Saccharomyces cerevisiae, Trichosporon asahii, Trichosporon inkin, Cryptococcus neoformans, Candida albicans, Candida tropicalis, Candida parapsilosis, Candida krusei, and...
and expressed in two-by-two contingency tables. The result for BAL fluid from treated versus untreated animals by Fisher’s exact test of the presence or absence of a positive GM EIA, qPCR, or quantitative culture survival and diagnostic BAL parameters was further analyzed by determination of water samples. No inhibition was observed in any of the samples. There was 80% sensitivity and 100% specificity at 36 cycles. Any crossover value ≤ 36 cycles was considered positive.

**RESULTS**

**Sensitivity and specificity of GM EIA, qPCR, and culture in BAL fluid.** For the GM assay the ROC showed 100% sensitivity and 100% specificity at GM index (GMI) cutoffs ranging from 0.75 to 4.0. A cutoff optical density ratio of 0.75 was chosen to define GM positivity for this study. Choosing a lower cutoff value would result in lower specificity while retaining sensitivity (100%). A higher cutoff value (above 4.0) results in decreased specificity while retaining sensitivity (Fig. 1A). The GM EIA of corresponding serum samples showed 100% sensitivity with an optical density cutoff ratio of 0.5.

The ROC for the qPCR assay indicated that a crossover value of 36 cycles would be the optimal point for positivity, resulting in a sensitivity of 80% and a specificity of 100%. Therefore, a cycle number ≤ 36 cycles was considered positive. A crossover value > 36 cycles resulted in increased sensitivity but decreased specificity and was therefore considered negative (Fig. 1B).

The sensitivity of BAL fluid culture as measured by log (CFU/milliliter) was 42% with a specificity of 100%. GM EIA was positive for 100% and qPCR was positive for 70% of the infected animals that BAL fluid culture failed to detect as infected (Table 1).

**Effect of antifungal therapy on sensitivity and specificity of GM EIA, qPCR, and culture of BAL fluid.** In analyses of the effect of antifungal treatment, different ROC results were observed for the GM EIA and qPCR assay. The GM EIA at our previously determined GMI cutoff of 0.75 had demonstrated a decrease in sensitivity to 92%, with the specificity remaining the same at 100% (Fig. 2A). This change corresponded to a 1% decline in the area under the ROC when antifungal therapy was administered. The sensitivity of the corresponding serum samples also declined with antifungal therapy to 90% (Table 1).

When antifungal therapy was administered, the sensitivity of the qPCR assay decreased to 50%; however, the specificity of the assay remained the same at 100% (Fig. 2B) while the ≤36 cycle criterion for positivity was retained. This decline in assay sensitivity corresponded to a 30% decline in the area under the ROC. However, in animals treated with antifungal therapy, the sensitivity of BAL fluid culture decreased even more to 16% (Table 1).

The GM EIA and qPCR assay of BAL fluid both showed a significant decline in signal for detection in experiments using rabbits treated with ravuconazole (P ≤ 0.02). In contrast, BAL...
fluid cultures did not show a significant decline in fungal burden with ravuconazole treatment ($P = 0.23$) (Table 2).

GM levels in BAL fluid remained elevated in rabbits treated with micafungin (Table 2), whereas BAL fluid from animals treated with micafungin showed a significant decline in signal when analyzed by qPCR ($P = 0.04$) (Table 2).

GM EIA, qPCR, and culture of BAL fluid demonstrated a treatment effect when deoxycholate amphotericin B was administered. All values were significantly lower than those of untreated controls ($P = 0.05$) (Table 2).

**Ability of BAL fluid to convey diagnostic or prognostic information.** In order to evaluate the relationship between qPCR analysis of BAL fluid and survival, we performed time sequence analysis and produced two-by-two contingency tables of diagnostic BAL parameters to address this question. Linear regression analysis of untreated controls demonstrated a trend towards an inverse relationship between survival ($y$) and qPCR result ($x$) [where $y = -0.26x + (7.3 \pm 0.54)$; $P = 0.067$]. Moreover, data with respect to the presence or absence of a positive qPCR result for BAL fluid demonstrated a significant difference ($P = 0.03$) between untreated controls (19/24; mean survival rate, 6.4 ± 0.3 days) and amphotericin B-treated animals (8/18; mean survival rate, 11.2 ± 0.3 days). Quantitative culture also revealed a significant difference ($P = 0.02$) between the results for untreated controls (11/24; mean survival rate, 6.4 ± 0.3 days) and amphotericin B-treated animals (2/18; mean survival rate, 11.2 ± 0.3 days).

**DISCUSSION**

The definitive diagnosis of IPA often requires invasive techniques such as biopsy of tissue for histology and culture (1). However, the diagnostic yield of conventional culture-based methods of BAL fluid may have a relatively low sensitivity for detection of *A. fumigatus* in patients with IPA (3, 19, 32). Incorporating diagnostic assays that are more sensitive and specific, such as GM EIA and qPCR, would permit earlier diagnosis of IPA, would improve clinical outcomes through timely initiation of antifungal therapy, and would allow withholding antifungal treatment when test results are negative. Additionally, both the GM EIA and qPCR assay may enable more rapid results than traditional culture-based methods (hours versus days).

In this well-established model of IPA, the ROC character-

![FIG. 2](http://jcm.asm.org/)

**FIG. 2.** (A) ROC demonstrating sensitivity and 1-specificity for the GM EIA at different GMI cutoff values for BAL fluid from 61 rabbits receiving antifungal therapy for IPA and 19 healthy rabbits. There was a decline in sensitivity to 92% with 100% specificity. (B) ROC demonstrating sensitivity and 1-specificity for qPCR at different cycle number cutoff values for BAL fluid from 61 rabbits receiving antifungal therapy for IPA and 19 healthy rabbits. There was a decline in sensitivity to 50%, with 100% specificity.
TABLE 2. Effect of antifungal therapy on detection of GM antigen, genomic DNA, and residual fungal burden in BAL fluid of animals with experimentally induced IPAa

<table>
<thead>
<tr>
<th>Treatment group (no. of animals)</th>
<th>EIA (GM)</th>
<th>PCR (log DNA/ml)</th>
<th>Culture (log CFU/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Untreated controls (24)</td>
<td>6.0 ± 0.18e</td>
<td>3.3 ± 0.46d</td>
<td>0.70 ± 0.22f</td>
</tr>
<tr>
<td>Animals treated with:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ravuconazole (17)</td>
<td>3.2 ± 0.50f</td>
<td>1.6 ± 0.47g</td>
<td>0.42 ± 0.24</td>
</tr>
<tr>
<td>Micafungin (26)</td>
<td>6.8 ± 0.30</td>
<td>2.1 ± 0.37h</td>
<td>0.21 ± 0.10i</td>
</tr>
<tr>
<td>Deoxycytidine amphotericin B (18)</td>
<td>4.7 ± 0.50i</td>
<td>1.2 ± 0.40j</td>
<td>0.16 ± 0.15k</td>
</tr>
</tbody>
</table>

a All P values (see below) represent comparisons of treatment groups to untreated controls.

b Data are means ± SEM.
c DNA data are expressed in femtograms.
d,e P < 0.0001.
e P < 0.05.
f P = 0.003.

g, h Not detectable.

ization of the GM EIA for BAL fluid had 100% sensitivity and specificity at an optimal GM cutoff of 0.75. These findings are consistent with those of Musher et al. (25), who found a similar ROC as well as optimal sensitivities and specificities between cutoff values 0.5 and 1.0. In other clinical studies (4, 34, 38), the GM assay had a sensitivity ranging from 85 to 100%. However, ROC analysis was not performed in these studies.

Due to documented cross-reactivity between GM antigen and antibiotics such as piperacillin-tazobactam, amoxicillin, and amoxicillin-clavulanic acid (2, 37, 39, 43), the specificity of the GM assay may be diminished in certain patient populations receiving these antibiotics. In these circumstances, the application of a second, more specific diagnostic test, such as qPCR, may provide important adjunctive information to improve diagnostic accuracy. The ROC characterization of the qPCR assay had a sensitivity of 80% and a specificity of 100% for BAL fluid. The threshold for detection of any extrinsic environmental contamination with conidia was ≥100 CFU.

Little is known about the effect of antifungal therapy on the release of GM and the detection of A. fumigatus DNA in BAL fluid. We therefore investigated the effects of antifungal treatment on the detection of GM antigen and A. fumigatus DNA in BAL fluid from neutropenic rabbits with experimentally induced IPA. ROC analysis performed with rabbits treated using antifungal therapy found lower sensitivities for GM EIA than untreated controls.

This study also found that antifungal therapy had a more profound effect on reducing the diagnostic yield of cultures than on that of GM EIA and qPCR. Antifungal therapy is widely used for prophylaxis or empirical therapy in high-risk patients. Consequently, culture-based assays of BAL from patients already receiving antifungal therapy may have diminished sensitivity. These findings are compatible with findings from a recent study of a cohort of patients with invasive aspergillosis; in that study, the sensitivity of GM EIA was diminished by previous antifungal therapy (23). Our findings indicate that the use of GM EIA and qPCR may complement culture-based methods and improve the analysis of BAL fluid for detection of A. fumigatus.

Triazole and echinocandin differed markedly in their effects on detection of GM antigen and genomic DNA of A. fumigatus in BAL fluid. Analysis of BAL fluid from rabbits treated with rauconazole demonstrated a significant decline in GM antigen as well as a decline in detectable signal of A. fumigatus DNA by qPCR. In contrast to the effects observed with rauconazole-treated rabbits, we found consistently elevated levels of GM with micafungin treatment. This paradoxical persistence in BAL GM levels in animals treated with micafungin may be due to the effect of the echinocandin in disrupting the cell wall and the inability of the drug to clear histologically detectable organism from lung tissue (30), thus causing the release of GM into the surrounding epithelial lining fluid of the alveoli.

To our knowledge this is the first study to fully characterize and compare GM EIA, qPCR, and quantitative culture for the detection of A. fumigatus in BAL fluid from a well-established animal model system of IPA. These data provide a scientific foundation for interpretation of data derived from clinical BAL fluid specimens from immunocompromised patients, for whom the diagnostic criteria may be less certain (1). The GM EIA and qPCR assay have greater sensitivity than culture in detection of A. fumigatus in BAL fluid. The performances of the GM EIA and qPCR assay may differ according to the specific type of antifungal therapy. These findings suggest that GM EIA and qPCR are useful tools in the analysis of BAL fluid for detection of A. fumigatus. Utilizing these assays in conjunction with culture-based diagnostic methods could facilitate an accurate diagnosis and more-timely initiation of targeted therapy and could result in fewer invasive procedures.

ACKNOWLEDGMENTS

We thank Heidi A. Murray and Christine Mya-San for their assistance in conducting laboratory animal studies. This research was supported by the Intramural Research Program of the National Institutes of Health and National Cancer Institute.

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