Evaluating A Semi-Nested PCR to Support Histopathology Reports of Fungal Rhinosinusitis in Formalin-Fixed Paraffin-Embedded Tissue Samples.

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Research Article

Keywords: Fungal rhinosinusitis, Formalin-fixed paraffin-embedded tissue, Histopathology, semi-nested PCR, Sequencing

DOI: https://doi.org/10.21203/rs.3.rs-558660/v1
Abstract

Background

Fungal rhinosinusitis (FRS) encompasses a various spectrum of disease, which vary in clinical presentation, histologic features, and biological significance. FRS is commonly classified in two categories, i.e., invasive and non-invasive infection. Histopathology is the “gold standard” diagnostic method, but it is not able to determine the genus and species. Almost more than 50% of the histopathologically proven cases, the culture elicited no reliable results. This study was an attempt to evaluate the diagnostic efficiency of semi-nested polymerase chain reaction (PCR) to confirmation of pathology reports obtained from formalin-fixed paraffin-embedded (FFPE) functional endoscopic sinus surgery (FESS) in FRS patients.

Methods

One hundred ten specimens were subjected to DNA extraction and histopathology examination. The amplification of β-globin gene by conventional PCR was used for confirming the quality of extracted DNA. The semi-nested PCR was performed using ITS 1 and ITS4 primers during two steps. In order to identification of causative agents, sequencing of the internal transcribed spacer region (ITS1-5.8S-ITS2) was performed on PCR products.

Results

Sixty-four out of 110 samples were positive by histopathology evidence, of which 56 samples (87.5%) were positive by PCR and 8 (12.5%) samples were negative. Out of 46 negative samples by histopathological methods, 41 samples (89.1%) were negative in PCR while 5 (10.9%) were positive. Sensitivity, specificity, positive predictive value, and negative predictive value of the semi-nested PCR method were reported 87.5%, 89.2%, 92.7%, and 85.2%, respectively. Aspergillus flavus, Rhizopus oryzae, Lichtheimia corymbifera and Candida albicans have identified as common fungal species.

Conclusion

Based on staining and direct examination that are time-consuming, applying molecular methods might be an appropriate choice for rapid diagnosis and support the consequences of histopathology examinations. We suggest that fresh biopsy specimens elicit more reliable results in comparison with FFPE samples. It is recommended that semi-nested PCR assays be performed in a single tube, which showed less prone to contamination when compared with assays that were carried out in two stages and in separate tubes.

Trial registration:

Not applicable.
Background

The inflammation of paranasal sinus mucosa is called sinusitis, or rhinosinusitis [1], which commonly affecting approximately 4.5 to 12% of the North America and European populations and 20% globally [2]. Direct costs of chronic rhinosinusitis estimated between 12 to 13 billion dollars per year in the United States [3, 4]. Depending on risk factors, colonization of fungal spores in sinonasal cavities triggers immunopathological consequences, fungal sinusitis, or more accurately fungal rhinosinusitis (FRS) [5]. FRS is being reported increasing worldwide frequency [5]. Elevation in the incidence rate of FRS is a consequence of global escalation in the immunocompromised population [6]. However, significant increasing of FRS reported in immunocompetent hosts without predisposing factors, the situation becomes more complicated [7].

FRS includes a wide range of clinical spectrum encompasses the mild form of superficial colonization and allergic manifestations to life-threatening invasive disease [5]. But histopathologically, FRS is divided into two categories, i.e., invasive and noninvasive infection, depending on tissue invasion of the mucosal layer. The invasive diseases include acute invasive (fulminant) FRS, granulomatous invasive FRS, and chronic invasive FRS, while, the noninvasive diseases include localized colonization, fungal ball, and fungus-related eosinophilic FRS that includes allergic fungal rhinosinusitis (AFRS) [5, 8, 9].

Diagnosis is always a big challenging in the management of FRS. In addition to clinical signs and symptoms and radiographic manifestations, which are non-specific, more reliable methods are available including direct microscopic examination by potassium hydroxide (KOH), histopathology by Gomori methenamine silver (GMS) and hematoxylin and eosin (H&E), culture methods by Sabouraud dextrose agar (SDA), Sabouraud-chloramphenicol (SC), and Sabouraud-chloramphenicol- Cycloheximide (SCC), antigen/antibody testing (Galactomannan and Alternaria antigen tests), and molecular methods [10, 11]. Moreover, the advanced molecular methods, single PCR, Nested PCR, semi-nested PCR, Quantitative Real-time PCR (qPCR), and hybridization with species-specific probes, are established [12–14]. Semi-nested PCR is a modified method planned to restrict non-specific binding in products due to the amplification of unexpected primer binding sites [12, 15]. The efficacy of molecular and mycological tests is indisputable. The culture of fungal agents of FRS may yield false positive or negative results respectively due to environmental contamination and loss of viability due to improper storage and transferring conditions [16]. Microscopic examination and histopathological techniques remain the gold standard for the detection of fungal elements in FFPE samples [17]. Moreover, histopathology plays a key role in the categorizing of this infection [11]. As the number of fungal elements in the specimens may be low, or not well stained, and the patient smear may not be examined by a trained professional, the test results may not have a proper sensitivity and specificity.

There are several limitations in collecting samples from FRS cases via endoscopy-guided biopsy, of which small amounts of obtained specimen that usually used only for histopathological diagnosis. Therefore, using a reliable molecular method for direct detection of FRS in FFPE samples might improve the diagnosis. Here, we evaluated the diagnostic accuracy and power of semi-nested PCR in comparison with the results of histopathology assay of FFPE in patients with FRS.
Materials And Methods

Samples & Patients

In the current prospective cross-sectional study, the FESS samples were collected from 110 patients (64 positive FRS and 46 non-FRS). The status of patients was proved by clinical sign and symptoms and computed tomography (CT) scan. Demographic data of the patients, including age, sex, type of operation, site of infection, background diseases, and final pathology report, were documented (Table 1). The paraffin blocks were prepared for cutting process by microtome. During this process, FESS samples were divided to 10 slices with 5 µm thickness. The samples were prepared and put into microtubes to further histopathological and molecular investigations.
Table 1
Demographic data of the patients and the results of molecular assays

| Patient No | Sex | Age | Risk Factors | Pathology Report | Outcome  | PCR β globin/ITS | Sequence          |
|------------|-----|-----|--------------|------------------|----------|-----------------|------------------|
| 1          | F   | 41  | ND           | Chronic FRS      | Survived | +/+             | A. flavus        |
| 2          | F   | 50  | ND           | Chronic FRS      | Survived | +/+             | C. albicans      |
| 3          | F   | 40  | ND           | Chronic FRS      | Survived | +/+             | A. flavus        |
| 4          | F   | 52  | Leukemia     | Mucormycosis     | Died     | +/+             | -                |
| 5          | F   | 42  | Diabetic     | Mucormycosis     | Survived | +/+             | R. oryzae        |
| 6          | F   | 50  | Leukemia     | Acute FRS        | Died     | +/+             | A. flavus        |
| 7          | M   | 55  | ND           | Mucormycosis     | Died     | +/-             | -                |
| 8          | M   | 2   | Leukemia     | Chronic FRS      | Survived | +/+             | A. flavus        |
| 9          | F   | 63  | Leukemia     | Mucormycosis     | Survived | +/+             | -                |
| 10         | M   | 21  | Allergy      | Chronic FRS      | Survived | +/+             | Cr. ozbekistanensis |
| 11         | M   | 51  | ND           | Chronic FRS      | Survived | +/+             | A. flavus        |
| 12         | M   | 26  | Leukemia - Allergy | Mucormycosis | Survived | +/-             | -                |
| 13         | M   | 58  | ND           | Chronic FRS      | Survived | +/+             | A. flavus        |
| 14         | F   | 47  | ND           | Chronic FRS      | Survived | +/+             | C. albicans      |
| 15         | M   | 32  | Allergy      | Chronic FRS      | Survived | +/+             | A. fumigatus      |
| 16         | M   | 26  | ND           | Chronic FRS      | Survived | +/+             | -                |
| 17         | F   | 42  | ND           | Mucormycosis     | Died     | +/+             | R. oryzae        |
| 18         | M   | 43  | ND           | Chronic FRS      | Survived | +/+             | -                |
| 19         | M   | 20  | Diabetic     | Mucormycosis     | Survived | +/+             | -                |
| 20         | F   | 49  | ND           | Mucormycosis     | Survived | +/+             | R. microsporus   |
| 21         | F   | 42  | Diabetic     | Acute FRS        | Survived | +/+             | A. oryzae        |
| 22         | M   | 25  | Leukemia     | Mucormycosis     | Survived | +/+             | -                |
| 23         | M   | 36  | Diabetic     | Mucormycosis     | Died     | +/+             | -                |
| 24         | F   | 44  | ND           | Mucormycosis     | Died     | +/+             | L. corymbifera   |
| 25         | F   | 53  | ND           | Mucormycosis     | Survived | +/+             | -                |
| Patient No | Sex | Age | Risk Factors          | Pathology Report | Outcome | PCR β globin/ITS | Sequence            |
|------------|-----|-----|-----------------------|------------------|---------|------------------|---------------------|
| 26         | F   | 35  | Leukemia              | Mucormycosis     | Died    | +/-              | -                   |
| 27         | F   | 32  | ND                    | Chronic FRS      | Survived | +/-              | A. alternate        |
| 28         | M   | 26  | ND                    | Chronic FRS      | Survived | +/-              | A. parasiticus      |
| 29         | F   | 70  | Leukemia              | Chronic FRS      | Survived | +/-              | A. flavus           |
| 30         | M   | 53  | Leukemia              | Mucormycosis     | Survived | +/-              | -                   |
| 31         | F   | 25  | Leukemia              | Mucormycosis     | Died     | +/-              | -                   |
| 32         | M   | 65  | ND                    | Mucormycosis     | Survived | +/-              | -                   |
| 33         | F   | 20  | Leukemia              | Mucormycosis     | Died     | +/-              | -                   |
| 34         | F   | 52  | Addiction             | Chronic FRS      | Died     | +/-              | A. flavus           |
| 35         | M   | 55  | ND                    | Mucormycosis     | Survived | +/-              | -                   |
| 36         | M   | 55  | Diabetic              | Mucormycosis     | Survived | +/-              | R. oryzae           |
| 37         | F   | 59  | Leukemia              | Mucormycosis     | Died     | +/-              | R. oryzae           |
| 38         | F   | 59  | Leukemia              | Mucormycosis     | Died     | +/-              | R. oryzae           |
| 39         | M   | 34  | Polyp                 | Acute and chronic FRS | Survived | +/-              | A. flavus           |
| 40         | F   | 37  | Leukemia              | Mucormycosis     | Died     | +/-              | -                   |
| 41         | F   | 64  | Autoimmune disease    | Mucormycosis     | Survived | +/-              | R. oryzae           |
| 42         | F   | 61  | ND                    | Chronic FRS      | Survived | +/-              | -                   |
| 43         | F   | 6   | Leukemia              | Mucormycosis     | Died     | +/-              | -                   |
| 44         | M   | 52  | Transplantation       | Mucormycosis     | Died     | +/-              | R. oryzae           |
| 45         | M   | 50  | Diabetic-Transplantation | Mucormycosis | Died     | +/-              | R. oryzae           |
| 46         | M   | 51  | Transplantation       | Mucormycosis     | Died     | +/-              | -                   |
| 47         | F   | 55  | Diabetic-Transplantation | Mucormycosis | Died     | +/-              | Saksenaea vasiformis |
| 48         | M   | 20  | ND                    | Mucormycosis     | Died     | +/-              | -                   |
| 49         | F   | 53  | ND                    | Mucormycosis     | Survived | +/-              | L. corymbifera      |
| 50         | F   | 51  | Diabetic-Leukemia     | Mucormycosis     | Survived | +/-              | -                   |
| Patient No | Sex | Age | Risk Factors | Pathology Report | Outcome | PCR β globin/ITS | Sequence |
|------------|-----|-----|--------------|------------------|---------|----------------|----------|
| 51         | F   | 52  | ND           | Chronic FRS      | Survived | +/+           | A. flavus |
| 52         | F   | 34  | Diabetic-Leukemia | Mucormycosis | Survived | +/+           | -        |
| 53         | M   | 56  | Diabetic     | Mucormycosis    | Died     | +/+           | -        |
| 54         | M   | 30  | Leukemia     | Mucormycosis    | Survived | +/+           | R. oryzae |
| 55         | F   | 62  | Diabetic     | Mucormycosis    | Died     | +/-           | -        |
| 56         | F   | 49  | Diabetic     | Chronic FRS     | Survived | +/-           | C. albicans |
| 57         | M   | 6   | Leukemia     | Mucormycosis    | Survived | +/-           | -        |
| 58         | F   | 46  | Diabetic     | Chronic FRS     | Survived | +/-           | A. flavus |
| 59         | F   | 22  | Leukemia     | Mucormycosis    | Died     | +/-           | -        |
| 60         | M   | 51  | Leukemia     | Mucormycosis    | Survived | +/-           | R. oryzae |
| 61         | M   | 33  | Diabetic     | Chronic FRS     | Survived | +/-           | A. flavus |
| 62         | F   | 47  | Leukemia     | Mucormycosis    | Died     | +/-           | -        |
| 63         | F   | 4   | Leukemia     | Mucormycosis    | Survived | +/-           | L. corymbifera |
| 64         | M   | 9   | Diabetic     | Mucormycosis    | Died     | +/-           | -        |

**Deparaffinization Process**

To reduce the contamination of samples during this process, it recommended the sterilization of microtome and other instruments using benzene and 2 M HCl rinsed with sterile water (www.leicabiosystems.com). The 1000 µl of xylene was added to microtube containing 5 µm of sample, which was then incubated in 56°C on a heating block for 15 min at room temperature and section subsequently centrifuged at 10,000 × g for 2 min. The supernatant was removed and 1,000 µl of absolute ethanol was added and followed by centrifugation at 10,000 × g for 3 min. The previous stage was repeated for 3 times then the tubes were incubated at 37°C on a heating block until the total evaporation of the ethanol [18].

**Histopathological assay**

The FFPE FESS samples were stained by hematoxylin and eosin (H&E), Periodic Acid Schiff (PAS) stains. The staining processes were performed according to the protocols for FFPE sample staining [19].

**DNA extraction**

DNA was extracted as previously described. Briley, 100 µl lysis buffer, 180 µl of ATL buffer, and 20 µl of proteinase K were added to the tube samples. After overnight incubation at 56°C, the tubes were washed via normal saline. To complete the lysis process, tubes were heated in boiling water for 5 minutes. The tubes
were incubated in boiling water and liquid nitrogen for 1 min and 2 min, respectively. This step was repeated several times. Finally, they reached room temperature. DNA extraction was completed by QIAamp DNA extraction from tissue mini kit (Qiagen, Hilden, Germany), as previously described [20]. This process is based on the binding of the DNA to silica columns.

**PCR Assay**

To evaluate the quality of the extracted DNA, human β-globin gene fragments amplified by the PCR method [21, 22] and KM29/PCO4 primers. PCR process was performed in a total volume of 25µl. So that, the master mix containing 1mM MgCl₂, 200 µM deoxyribonucleotide triphosphates solution (dNTP)s, 1X reaction buffer10x, 1U Taq DNA polymerase (total of Cinna Gene, Iran), and 1µl (10 picomols) of forward and reverse primers (Table 2). PCR conditions were as follows; denaturation phase 1 cycle at 94°C for 10 min followed by 35 cycles of 94°C for 45 s, 58°C for 45 s and 72°C for 5 min, and a final extension at 72°C for 7 min. Amplicon quality and concentrations were estimated on the agarose gel and analyzed by the Gel Doc XR system (Biorad, USA). The smart Ladder (Eurogentec, Seraing-Belgium) was used as the size and concentration marker.

| Primers          | 5 → 3 sequence    | References |
|------------------|-------------------|------------|
| β globin (F:KM29)| GGT TGG CCA ATC TAC TCC CAG G |            |
| β globin (R:PCO4)| CAA CTT CAT CCA CGT TCA CC   |            |
| ITS1 (F)         | TCC GTA GGT GAA CCT GCG G    |            |
| ITS2 (R)         | GCT GCG TTC TTC ATC GAT GC   |            |
| ITS4 (R)         | TCC TCC GCT TAT TGA TAT GC   |            |

**Semi-nested PCR Assay**

Universal fungal ITS region (ITS1-5.8S-ITS2) was targeted for evaluation by semi-nested PCR. The first PCR reaction was performed using ITS1 (forward) and ITS4 (reverse) primers (Table 2). The total volume was 50µl encompasses 45 µl of reaction mixture containing 1 mM MgCl₂, 1x of PCR buffer 10x, 0.1 mM each deoxynucleotide triphosphate (dNTP), 0.5 pmols/µl of each primer, 1 U of Taq DNA polymerase, and 5 µl of nucleic acid extract. In the second PCR step, the ITS1 (forward) and ITS2 (reverse) regions were amplified within the 3 µl of diluted (1/100) product of the first PCR step. The product of first step was run into the second PCR to amplify ITS1 (forward) and ITS2 (reverse) regions in the same total volume (Table 2). PCR conditions for the first step were as follows; 10 min of initial denaturation at 94°C, 35 cycles of 96°C for 45 s, 58°C for 45 s and 72°C for 1 min, and a 5 min final extension at 72°C. Also, PCR conditions for the second step were as follows; 5 min of initial denaturation at 94°C, 32 cycles of 94°C for 30 s, 60°C for 30 s and 72°C for 45 s, and 5 min final extension at 72°C [15]. Both steps of the semi-nested PCR encompassed 10 to 20 samples included a positive control containing 0.5 ng of purified DNA of one of the fungal isolates and at
least two blanks with reagents only. Product quality and concentrations were estimated on the agarose gel, and analyzed by the Gel Doc XR system (Biorad, USA). Smart Ladder (Eurogentec, Seraing, Belgium) was used as the size and concentration marker.

**Sequencing**

To further confirmation of the results, PCR products were bidirectional chain terminated Sanger sequencing using referenced primers was performed. The obtained sequences were searched using the NBLAST algorithm (https://blast.ncbi.nlm.nih.gov/Blast.cgi) and the identity of each strain was assigned accordingly.

**Data Statistics**

Data were analyzed using SPSS software version 24. Briefly, descriptive data were presented as mean, standard deviation, percentages, and charts. The Chi-square and Fisher's exact tests were used to compare qualitative variables between the two groups. The Student t-test was used to compare quantitative variables between the two groups. Also, the Agreement between the two diagnostic methods was calculated by Kappa test, interpretation of Kappa was based on Viera et al. [23]. The p-value less than 0.05 (P < 0.05) was considered statistically significant.

**Results**

**Patients and samples**

One hundred ten FFPE samples were retrospectively collected from patients for two years (2018 to 2020). Briefly, 50 patients were male (45.5%) between 2 to 82 years old (mean age: 40.2 years old). The demographical data of the patients were truncated in the Table 1. Most patients with FRS had predisposing factors, which respectively including leukemia (20:64, 31.23%), diabetes (11:64, 17.11%), transplantation (4:64, 6.25%), allergy (3:64, 4.7%), polyp, autoimmune diseases, and drug addiction (1:64, 1.56%). The crude mortality rate among 64 patients with FRS was 37.5% (24/64) but the mortality rate in mucormycosis was 53.7% (22/41).

**Histopathological Examinations**

Sixty-four samples were proven to positive for FRS. In 64% (41 of 64) of them, ribbon-like non-septate or minimally septate hyaline mycelium were seen indicating mucormycosis (Figs. 1–3). Moreover, pathology evidence indicated that 31% (20 of 64) of FRS cases were chronic FRS, 3.12% (2 of 64) were acute, and 1.56% (1 of 64) was acute/chronic FRS (Table 1). Forty-six non-FRS samples were included as controls.

**Molecular assay**

As shown in Table 3, 56 out of the 64 (87.5%) histological proven specimens were positive for PCR (as shown in Fig. 4), and 8 samples (12.5%) were negative. Out of 46 negative samples by histopathological methods, 41 samples (89.1%) remained negative for PCR, while 5 (10.9) were positive. Moreover, 100% (23/23) of chronic, acute, and acute/chronic FRS cases and 80.5% (33/41) of mucormycosis cases were positive via semi-nested PCR method. Sensitivity, specificity, positive predictive value and negative predictive values of semi-nested PCR were 87.5%, 89.1%, 92.7%, and 85.2%, respectively. Moreover, kappa factor
between these two methods were 0.76 indicating substantial agreements between these two tests. From the 56 samples that were sequenced, 35 PCR product were successfully identified. The thirty-five sequences generated in this study were deposited in the GenBank database (accession numbers from MZ333236 to MZ333270 with persistent accessible links form MZ333236 to MZ333270, respectively). Eventually, \textit{Aspergillus flavus} (12/35), \textit{Rhizopus oryzae} (10/35), \textit{Lichtheimia corymbifera} (3/35), \textit{Candida albicans} (3/35) \textit{Aspergillus oryzae} (1/35), \textit{A. parasiticus} (1/35), \textit{A. fumigatus} (1/35) \textit{Saksenaea vasiformis} (1/35), \textit{Rhizopus microsporus} (1/35), \textit{Alternaria alternata} (1/35), and \textit{Cryptococcus uzbekistaniensis} (1/35) were identified by sequencing of the ITS region.

| Pathology report | PCR result | Number (%) |
|------------------|------------|------------|
| FRS (n = 64)     | Positive   | 56 (87.5)  |
|                  | Negative   | 8 (12.5)   |
| non-FRS (n = 46) | Positive   | 5 (10.9)   |
|                  | Negative   | 41 (89.1)  |

**Table 3**
Comparing the results of histopathology and PCR assay

**Discussion**

By scrutiny in-field studies, several factors found to be affected the consequence of PCR assay in FFPE tissue samples. These factors are including DNA extraction method, inclusion of proper housekeeping gene, PCR method (i.e., panfungal, specific, nested, semi-nested, multiplex, and Real-Time), target gene(s) primers, amplicon length, thickness of FFPE cut, storage time of specimen, and contamination during sample preparation [24–26].

Here, we reached a sensitivity of 87.5% and a specificity of 89.1% for semi-nested PCR assay targeting ITS1-5.8s-ITS2 (ITS1-2) region of 110 FFPE-FESS samples. Bialek \textit{et al.} indicated a 60.6% sensitivity and 100% specificity for nested PCR targeting 100-kDa-like-protein gene in 33 FFPE specimens, which were histopathologically proven for histoplasmosis [27]. Similarly, Willinger \textit{et al.} indicated the 87% sensitivity for PCR of the 28S region [28]. This may result from short amplicon length (260 bp) and pertinent primer pairs (U1/U2) to amplification the 28S region. In another similar study, Rickerts \textit{et al.} reported acceptable sensitively for molecular diagnosis of invasive aspergillosis and mucormycosis [29]. In both studies [22, 29], the DNA extraction method, gene targets, primer pairs, and amplicon length were similar. Lau \textit{et al.} reported 82% positive results for panfungal PCR assay [30]. They also showed a sensitivity of 97% and 68% for fresh and FFPE samples, respectively which was lower than the results of current study. Cabaret \textit{et al.} indicated
62.5% and 93.75 positive results of FFPE samples for conventional PCR and qPCR, respectively [31]. These differences may result from the differences in amplicon length (> 300 bp for conventional PCR vs. 150 bp for qPCR) and the type of PCR method. In a similar study, Hammond et al. reported 81.5% positive results by semi-nested PCR of 18S rDNA using ZM1/ZM2 and ZM1/ZM3 primer pairs targeting an amplicon < 200 bp [32]. Salehi et al. reported a 64% sensitivity for identification of fungi from FFPE by qPCR assay [33]. Drogari-Apiranthitou et al. [34] reached 45% positive PCR for *Mucorales* and 40% positive for *Aspergillus* spp. They reported 79.3% and 100% sensitivity and specificity for semi-nested PCR, respectively. The higher sensitivity of their method in comparison to us might be due to the more thickness of their tissue cuts (10 µm vs. 5 µm).

In another study, Ganesan et al. reported the site of infection might affect the sensitivity of PCR test and reported that sensitivity rate increased from 63–83% for angioinvasion sites [35]. While Jung et al. indicated a 41.3% positive results for panfungal PCR, due to long amplicon length and may be inappropriate DNA extraction kit [36]. In our study we found a concordance rate of 76% for semi-nested PCR of the ITS region and histopathology assays which is similar to previous reports [30]. While others reported higher [33] or lower [37, 38] concordance rate between these two methods.

Due to the low amount of tissue in FFPE samples, the extracted DNA would be low. Hence, selection of proper extraction method is a key step in achieving a reliable amplification. Here, we used QIAamp DNA FFPE Tissue Kit (QIAGEN), as successfully used before [17, 28, 34, 37–43]. In a comparative study between different commercial extraction kits, Muñoz-Cadavid et al. reported TaKaRa and QIAamp extraction kits yielded the best results for extraction of high quality DNA from the FFPE sample [44].

Similar to some previous studies [22, 32–34, 41, 44], we used human β-globin gene as internal control and the primers successfully amplified the expected region in all samples. While others used different genes such as IRBP [38], GAPDH [27], and ZP3 [45] as controls.

In this study we used a panfungal semi-nested PCR for amplification of the ITS1-5.8s-ITS2 followed by ITS1 regions to detect fungal element in FFPE tissue samples which is concordant with a previous report [24]. Others amplified different targets such as 28S rDNA [43], ITS2 [35], 18S rDNA [34, 37] and mitochondrial tRNA [34] for detection of fungal DNA in FFPE samples. Although, Cabaret et al. suggest targeting mitochondrial DNA is superior to ITS [31], they obtained lower positive rate by conventional PCR (10/16, 62.5%) in comparison to ours (56/64, 87.5%). In an another study, Jillwin et al. [37] targeted five different gene regions including universal ITS (ITS1-5.8s-ITS2), ITS1, ITS2, 18S rDNA, and D1/D2 of 28S rDNA and reported that ITS1 amplification leads to 61.9% positive results by PCR method.

The false-negative/positive consequences may result from the following reasons: First, false-positive histopathological results artifacts during staining. Second, the presence of conserved genes in multiple copies (rDNA) is a disadvantage in clinical specimens collected from nonsterile body sites, because nonpathogenic commensal fungi, environmental spores, or colonizing fungi can also cause considerable nonspecific amplification in samples consisting mainly of human cells with only a few fungal cells. Third, proteins-DNA cross-linking formation and fragmentation of DNA during the fixation process leads to a lack of intact DNA needed for amplification. More specifically, it is difficult to amplify the target gene when DNA is highly fragmented or cross-linked and the amplicon size is large. Fourth, amplification of a housekeeping
gene (human β-globin) in tissue does not necessarily mean the presence of a sufficient amount of amplifiable fungal DNA.

It is recommended that a shorter amplicon length gives higher sensitivity and specificity rates for PCR results [26]. Also, it recommended that semi-nested PCR assays performed in a single tube, which showed less prone to contamination when compared with assays that carried out in two stages and in separate tubes. As *Aspergillus* species and Mucorales are most frequent causative agents of FRS, it can be a good idea to use species-specific primers and multiplex PCR to differentiate them as soon as possible in a single reaction.

**Conclusion**

In this study amplification of ITS1-5.8S-ITS2 by semi-nested PCR in FFPE samples of IFI yielded a considerable result with sensitivity and specificity of 87.5% and 89.1%, respectively. Considering low amounts of samples obtained by endoscopy-guided biopsy and rapid clinical progress of rhinocerebral mucormycosis, establishment a rapid, accurate and reliable molecular method for direct detection of FRS in FFPE samples might improve the outcomes.

**List Of Abbreviations**

FRS  
Fungal rhinosinusitis  
PCR  
polymerase chain reaction  
FFPE  
formalin-fixed paraffin-embedded  
FESS  
functional endoscopic sinus surgery  
ITS  
internal transcribed spacer  
AFRS  
allergic fungal rhinosinusitis  
KOH  
potassium hydroxide  
GMS  
Gomori methenamine silver  
H&E  
hematoxylin and eosin  
SDA  
Sabouraud dextrose agar  
SC  
Sabouraud-chloramphenicol  
SCC
Declarations

Ethics approval and consent to participate: All methods were performed in accordance with the relevant guidelines and regulations. This study were supervised and monitored by the Ethics Committee of Shiraz University of Medical Sciences (permission code: IR.SUMS.REC.1393.S7348). Informed consent were taken from all participants and parents/legally authorized representative of all minor (age less than 16 years) and deceased participants involved in the study.

Consent for publication: Not applicable.

Availability of data and materials: The thirty-five sequences generated in this study were deposited in the GenBank database (accession numbers from MZ333236 to MZ333270 with persistent accessible links form MZ333236 to MZ333270, respectively). All data used to support the findings of this study are available from corresponding author upon request.

Competing interests: The authors declare that they have no competing interests.

Funding: This project received financial support from the Vice President of Research, Shiraz University of Medical Sciences (Grant & Thesis No. 937348).

Authors' contributions: M. JA, M.K, K.P, H.M and K.Z designed the study. M.JA, G.S, B.K, and M.S collected and prepared study samples. M.K, S.A, K.A, F.S, and K.Z performed molecular assays and data analysis. M. JA, G.S, performed histopathological tests. M.K, H.M, H.B, K.P, and K.Z drafted the manuscript, acquired the data; analysed and interpreted the data. All authors read, revised and approved the final manuscript.

Acknowledgements: The authors are grateful to Khalili Educational and Research Hospital, Shiraz-Iran and Vice President of Research, Shiraz University of Medical Sciences for great support of this project.

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**Figures**
Figure 1

Mucor hyphae in a necrotic background of sinusal tissue, H&E stain (x400)

Figure 2

Fungal spores in the sinusal tissue of chronic sinusitis, H&E stain (X400)
Figure 3

non-septate hyphae in the sinusal tissue, H&E stain (x400)

Figure 4

Gel electrophoresis primary PCR via β-globin (left) and semi-nested PCR for ITS region (wright)