A Whole Genome Sequencing-Based Approach to Track down Genomic Variants in Itraconazole-Resistant Species of Aspergillus from Iran

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Abstract: The antifungal resistance in non-fumigatus Aspergillus spp., as well as Aspergillus fumigatus, poses a major therapeutic challenge which affects the entire healthcare community. Mutation occurrence of cyp51 gene paralogs is the major cause of azole resistance in Aspergillus spp. To obtain a full map of genomic changes, an accurate scan of the entire length of the Aspergillus genome is necessary. In this study, using whole genome sequencing (WGS) technique, we evaluated the mutation in cyp51A, cyp51B, Cdr1B, AtrR, Hmg1, HapE and FfmA genes in different clinical isolates of Aspergillus fumigatus, Aspergillus niger, Aspergillus tubingensis, Aspergillus weillwitschiae and Aspergillus terreus which responded to minimum inhibitory concentrations of itraconazole above 16 µg mL−1. We found different nonsynonymous mutations in the cyp51A, cyp51B, Cdr1B, AtrR, Hmg1, HapE and FfmA gene loci. According to our findings, Aspergillus species isolated from different parts of the world may represent different pattern of resistance mechanisms which may be revealed by WGS.

Keywords: Aspergillus spp.; itraconazole resistance; cyp51 genes; whole genome sequencing

1. Introduction

The mycoses caused by Aspergillus spp. include a wide variety of pathophysiological disorders, ranging from allergic manifestations to systemic infections [1]. Among the invasive fungal diseases (IFD), invasive aspergillosis (IA) is one of the most severe clinical entities, with mortality rates up to 70% in more than 300,000 cases estimated worldwide [2]. Although several studies have been focused on the most common species of Aspergillus including A. fumigatus and A. flavus, using molecular taxonomic tools, cryptic/rare Aspergillus species have recently been identified [3]. Hence, there is a growing awareness about the clinical relevance of Aspergillus cryptic species in the development of IA [4,5]. According to clinical observations, the epidemiological distribution of cryptic Aspergillus species are associated with invasive infections and often falls into the sections Fumigati, Flavi, Nigri, and Terrei [6]. Azole derivatives are regarded as the first-line options for the prophylactic treatment of IA [7]. Among azole antifungals, itraconazole (ITC) is widely used in medicine and veterinary treatment. This antifungal agent was considered as the treatment of choice for chronic pulmonary aspergillosis [8], and have been proposed as an...
alternative treatment for allergic bronchopulmonary aspergillosis and the management of patients with severe asthma [9]. An unsettled dilemma in managing aspergillosis crisis is the emergence of azole-resistance in *Aspergillus* spp., a phenomenon that stems from intricate mechanisms [10]. Due to the global rampancy of *A. fumigatus*, major studies have been steered to scrutinize the mechanisms and understand the roots of resistance against triazoles derivatives including ITC in environmental and clinical isolates [11–13]. With trailing the function of genes, it is inferred that occurrence of point mutations in *cyp51* gene paralogs (*cyp51A* and *cyp51B*) in *A. fumigatus*, *A. niger* and *A. terreus* are the most common genomic changes leading to azole resistance [14]. In response to triazoles acting on the locus of *cyp51A* and *cyp51B* genes, resulting mutation manifests as a non-synonymous substitution that alters the sequence of encoded proteins and thus resistant mutants are aroused [15,16]. However, mutation in non-*cyp51* gene including ATP-binding cassette transporters (*Cdr1B* or *abcG1*), *AtrR*, *Hmg1*, *HapE* and *FfmA* gene loci have been recently reported [17–21]. In recent years, whole genome sequencing (WGS) and bioinformatics methodologies have gained traction in the diagnostic field as potent appliances for detecting antifungal resistance in medical mycology [22]. Using WGS with a high number of clinical and environmental isolates of *A. fumigatus*, Rhodes et al. [23] showed that a strong genetic structuring into different clades with little interclade recombination, high confidence documents to show that transmission of resistant isolates occurs from the environment to the infected patients. The selective sweeps across multiple regions indicate a polygenic basis to traits in some genetic backgrounds, and the genes related to the azole resistance with previously unknown resistance mechanisms. Since WGS is a high-throughput technology that reads out the whole genomic content, it provides high-resolution images of single nucleotide polymorphisms (SNPs) at different points in the DNA of the target microorganism, which allows for wider genomic analysis as compared to primer design-based techniques [24]. In light of lack of information on WGS related to azole-resistant *Aspergillus* spp. and the importance of clinically relevant and cryptic species of *Aspergillus*, we evaluated the function of WGS to detect the genomic variations of *cyp51A*, *cyp51B*, *Cdr1B*, *AtrR*, *Hmg1*, *HapE* and *FfmA* genes in ITC-resistant isolates of *A. fumigatus* (section *Fumigati*), *A. terreus* (section *Terrei*), *A. niger*, *A. tubingensis*, and *A. welwitschiae* (section *Nigri*).

2. Materials and Methods

2.1. Clinical Isolates

A total of seven identified (using molecular techniques by sequencing of the β-tubulin-encoding gene (*benA*)) *Aspergillus* isolates including *A. fumigatus* (AF), *A. niger* (AN) and *A. welwitschiae* (AW), and *A. tubingensis* (AT1 and AT2) and *A. terreus* (AR1 and AR2) that responded to minimum inhibitory concentrations (MICs) of ITC above 16 µg mL⁻¹ were included in the study. Table 1 shows the clinical source and accession number of isolates.

| Sample Code | Species       | Section | Accession Number (benA Gene) | Origin                        |
|-------------|---------------|---------|------------------------------|-------------------------------|
| AF          | *A. fumigatus*| Fumigati| MH208797                     | Patients with ABPA           |
| AN          | *A. niger*    | Nigri   | MH208734                     | Nail (Onychomycosis)         |
| AW          | *A. welwitschiae* | Nigri   | MH208730                     | Sputum (asthmatic patient)   |
| AT1         | *A. tubingensis* | Nigri   | MH208733                     | BAL (Patient with probable IA) |
| AT2         | *A. tubingensis* | Nigri   | MH208779                     | Nail (Onychomycosis)         |
| AR1         | *A. terreus*  | Terri   | MH208803                     | Sputum (asthmatic patient)   |
| AR2         | *A. terreus*  | Terri   | MH208818                     | BAL (Patient with probable IA) |

**Table 1.** The clinical source and accession number of isolates.

BAL: Bronchoalveolar lavage fluid, IA: invasive aspergillosis, ABPA: Allergic Bronchopulmonary aspergillosis.

2.2. Molecular Identification

DNA was extracted from fungal colonies grown for 5–7 days on malt extract agar (Difco, Detroit, MI, USA) using a previously described method [25]. Polymerase chain
reaction (PCR) for partial amplification of the \textit{benA} was performed using forward and reverse primers Bt2a (5′-GGTAACAAAATCGGTGCTGCTTTC-3′) and Bt2b (5′-ACCCTCAGTGTAGTGACCCCTGGC-3′), respectively [26]. PCR products were analyzed by gel electrophoresis using 1.5% agarose gels, and the amplified \textit{benA} fragments were sequenced using the same primers with the ABI3730XL Genetic Analyzer platform (Applied Biosystems, Waltham, MA, USA). All sequences were subjected to nucleotide BLAST search of the NCBI database and the molecular database of the WI-KNAW Fungal Biodiversity Center (Utrecht, The Netherlands), and isolates were identified up to the species level.

2.3. Antifungal Susceptibility Testing

Antifungal susceptibility test (AFST) was fulfilled as a 2× microdilution broth method based on the Clinical and Laboratory standards Institute (CLSI)-M38-A3 instructions [27]. The azole antifungals evaluated in this study were ITC, voriconazole (VOR), posaconazole (POS), and isavuconazole (ISV) that purchased from Merck Sharp and Dohme BV, Haarlem, The Netherlands. To test antifungal susceptibility, a suspension of each isolate was prepared in RPMI 1640 broth medium (absorbance reading, A\textsubscript{530} of 0.09 to 0.13). Five azole derivatives mentioned above were also diluted in dimethyl sulfoxide (DMSO). \textit{Aspergillus} suspensions in 96-well plates were exposed to azoles by 2× dilution (ranging from 0.016 to 16 µg mL\textsuperscript{−1}) and incubated at 35 °C for 48 h. \textit{Candida parapsilosis} (ATCC 22019) and \textit{Pichia kudriavzevii} (ATCC 6258) were used as quality control strains. The MIC of the isolates was evaluated and after comparing with defined ECV values in CLSI supplement M59, non-wild type (NWT) isolates were identified.

2.4. Workflow of Whole Genome Sequencing (WGS)

DNA extraction was performed on each isolate after a three-day culture with the DNA extraction kit (SimBiolab, Mashhad, Iran). Genomic DNAs were quantified and assessed qualitatively using gel electrophoresis and NanoDrop spectrophotometer. The extracted gDNAs were transferred to Novogene (Tianjin, China) for genomic sequencing using GenTegra-DNA tubes (USA), which protect against degradation. Prior to starting the sequencing process, an RNAase treatment step was also considered to improve gDNAs purity. The DNAs were subjected to a 5400 fragment analyzer system, designed to meet the needs of ensuring their quantity, integrity and purity, as the preliminary assaying types. The libraries were prepared at Novogene Bioinformatics Technology Co., Ltd. (Beijing, China) using TruSeq\textsuperscript{®} DNA PCR-Free Sample Preparation Kit (Illumina, San Diego, CA, USA) according to manufacturer’s recommendation. In brief the genomic DNA was randomly sheared into short fragments. Then, the obtained fragments were end repaired, A-tailed and further ligated with Illumina adapter. The fragments with adapters were PCR amplified, size selected, and purified. The quality of the prepared libraries was assessed using Qubit and real-time PCR. At that time, a library was prepared according to each of the targeted isolates. After going through all these steps, all DNA samples were sequenced using paired-end 150 based pair sequencing. As part of our whole-genome sequencing project, we used an Illumina NovaSeq 6000 system, and the sequencing depth for the isolates was 100×. To conclude, data generated by Illumina platform were processed via CLC genomics workbench 21 software (QIAGEN GmbH, Hilden, Germany) according to following steps:

\begin{itemize}
  \item[i.] The forward and reverse sequencing data were imported into the software using Illumina High-Throughput Sequencing import option (Illumina Pipeline 1.8) in which the data were paired-ended, de-multiplexed and trimmed. The failed reads were also discarded.
  \item[ii.] The quality of the sequencing data was assessed using quality control of the software which include a complete list of lengths and quality distribution, GC content, ambiguous base-content, nucleotide contributions, sequence duplication levels and the sequence of the duplicated reads.
  \item[iii.] The reference sequence of each species was downloaded from NCBI (Table 2) and used as a reference for data mapping and resequencing using the Map Reads to Reference
option of the software, then the quality of the mappings was assessed using QC for
Read mapping and Genome Coverage Analysis option of the software.
iv. The variant calling of the assembled sequencing data was performed using Basic Variant
Detection option of the software, then each variant was also annotated manually
of the target genes for confirmation.
v. Finally, the functional consequences of the variations was determined using Amino
Acid Change option of the software by comparison of the variant tracks of the assem-
bled sequence with the references.

All DNA sequence data were uploaded to NCBI SRA-submission portal (https://
submit.ncbi.nlm.nih.gov/subs/sra/ (accessed on 20 November 2021)). In Figure 1, the
steps from the research are depicted.

Figure 1. The steps of project implementation, from sample collection to WGS. The estimated time to
complete each of the key stages of the project indicates that after the sampling phase, which spans
a period of one year, within two months, it is possible to analyze azole-resistant Aspergillus spp.
through whole genome sequencing to identify all SNPs in the entire genomic content.

2.5. Homology Modeling and Mutation Mappings

The primary amino acid sequence for CYP51A of A. fumigatus (XP_752137.1), A. wel-
witschiae (XP_026629604.1), Aspergillus niger (XP_001394224.1), Aspergillus tubingensis
(XP_035357335.1) and Aspergillus terreus (XP_001215095.1) were used to find appropriate
template for homology modeling of the 3D structure of each protein using ICM-Pro soft-
wire (Molsoft L.L.C., San Diego, CA, USA). Each template was imported into the software,
refined and minimized then used for homology modeling as described previously [28].
3. Results
3.1. Determination of MIC

The MIC values for seven isolates tested with ITC, VOR, POS, and ISV are provided in chart 1. Due to the presence of an ECV definition in CLSI M59 supplement for *A. fumigatus*, *A. niger*, and *A. terreus*, the MIC profiles of the four isolates AF, AN, AR1, and AR2 were comparable. AF, AN, AT1 and AT2 response to ITC with MIC > 16, POS = 1, VOR = 2, 1, 1 and 0.5, ISV = 1, 2, 0.5 and 0.5, respectively. The four isolates with MIC > 16 to ITC were therefore identified as NWT isolates (Figure 2A). For the three isolates AW, AT1 and AT2, there was no standard ECV to comparison. AW, AT1 and AT2 isolates showed MIC > 16 to ITC, MIC = 1 to POS, MIC = 2, 2 and 0.25 to VOR and MIC = 2, 0.5 and 1 to ISV, respectively (Figure 2B).

![Figure 2](image-url)

*Figure 2.* (A) The distribution of minimum inhibitory concentrations (MICs) of different *Aspergillus* isolates against tested triazoles compared to epidemiological Cutoff values (ECVs). (B) The MICs of different *Aspergillus* isolates without defined ECV against tested triazoles.
The detailed information on the isolates is also accessible in our recent published paper [3].

3.2. DNA Quality Control (QC) Report

During quality control, all seven isolates passed the criteria for integrity, concentration, and purity. DNA concentrations were between 50 and 75 ng/µL, and integrity tests yielded satisfactory upshots.

3.3. Homology Modeling and Mutation Mappings

Conformational structure of A. welwitschiae was not shown as the V16L mutation is located on the signal peptide of CYP51A.

In the current study, different mutations were identified and mapped on the structure of the modeled CYP51A. As illustrated in Figure 3, some mutations were located near the channel entry especially in A. Fumigatus (Figure 3D) but far from the active site of the proteins. This could provide a potential mechanism for azole resistance by limiting the access of inhibitors to the active site.

![Figure 3](image-url)

Figure 3. Homology modeling of CYP51A of the selected Aspergillus species and mutation mapping on the structures. Each structure was shown in two different presentations. (A) (A. terreus), (B) (A. tubingensis), (C) (A. niger) (D) (A. fumigatus).

3.4. Bioinformatics Analysis

Upon alignment of the sequences of cyp51A, cyp51B, Cdr1B, AtrR, Hmg1, HapE and FfmA as target genes confines of the seven isolates with the reference sequences deposited in NCBI, genomic variations were detected at different positions, which are depicted in Table 2. The intron sequences have been removed in the raw data processing process. The SNP positions in Table 2 are based on DNA coding sequences (CDS) that corresponds to the sequence of amino acids in a protein. Four nonsynonymous mutations have been identified in cyp51A sequence (T137A (F/Y), A514G (M/V), A743T (N/T) and C765G (D/E)) of A. niger isolate. Although the occurrence of the other four SNPs did not start off amino acid changes (G267A, A1047G, G1279A and T1362C). A. niger isolate contained three nonsynonymous substitutions (A169G (T/A), A683G (Q/R) and G1147C (V/L)) and eleven silent substitutions across the cyp51A gene. In cyp51B gene, variation at points A800T (N/I) and A1184C (N/T) altered the protein sequence. A solitary A. welwitschiae (AW) isolate was consorted with one and three replacement mutations in cyp51A (G46C (V/L)) and cyp51B genes (A440G (E/G), T525A (N/K), and A2785G (I/V)), respectively. Two isolates of A. tubingensis with AT1 and AT2 codes revealed dissimilar genomic changes in
the cyp51A and cyp51B genes. The former showed only one nonsynonymous mutation in the cyp51A target (C166T (L/F)) and the latter showed the same pattern as the reference genome in the cyp51A target. The amino acid sequences of isolates AT1 and AT2 were unaffected by point mutations in the cyp51A and cyp51B genetic codes. A. terreus isolates also gave different results. For AR1, replacement mutations at C662T (A/V) of the cyp51A gene, and at T1166G (T/S) of the cyp51B gene were unmask. In the second isolate of A. terreus, AR2, a silent mutation was spotted in the cyp51A gene, while no meaningful variance was stood up along the cyp51A gene as compared to reference sequence. It has been found that all seven AN, AF, AW, AT1, AT2, AR1 and AR2 isolates ultimately showed resistance to ITC following occurrence of substitution mutations at different loci of cyp51A and cyp51B and as target genes under study (Table 2). In non-cyp51 gene loci, nonsynonymous mutations have been identified in different isolates as follows:

A. weitschiae in Cdr1B (one mutation), Hmg1 (five mutations) and HapE (one mutation). A. fumigatus in AtrR and Hmg1 (two mutations, each). A. niger in Cdr1B (four mutations), AtrR (two mutations) and Hmg1 (one mutation). Isolate AT1 of A. tubingensis in Cdr1B (tbow mutations), HapE (one mutation) and FfmA (two mutations). Isolate AT2 in AtrR and FfmA (two mutations, each).

Isolate AR1 of A. terreus showed one mutation in HapE locus and AR2 in Cdr1B (one mutation), and AtrR (two mutations) (Table 2).

Our data is associated with an NCBI BioProject accession, PRJNA845900.

Table 2. The list of nonsynonymous substitutions along different gene loci in study Aspergillus isolates.
Table 2. Cont.

| Species | BioSample Accessions | Cyp51A | Cyp51B | Cdr1B | AtrR | Hmg1 | HapE | FfmA | Reference Accession Number |
|---------|----------------------|--------|--------|--------|------|------|------|------|---------------------------|
| AT2     | SAMN28864582         | -      | -      | -      | -    | -    | -    | T1293C(S/P) T1468C(F/S)   | GeneID:56006014 GeneID:56008603 GeneID:56003599 GeneID:56007043 GeneID:56006955 GeneID:56001387 GeneID:56005349 |
| AR1     | SAMN28864583         | C706T (A/V) | C1280G (T/S) | -      | -    | -    | T1103C (F/S)   | GeneID:4321797 GeneID:4317693 GeneID:4319285 GeneID:4318923 GeneID:4354193 GeneID:4321255 GeneID:4315909 |
| AR2     | SAMN28864584         | -      | -      | A3239T (E/D) | A70C (N/H) A2735C (Q/H) | -    | -    | -    | GeneID:4321797 GeneID:4317693 GeneID:4319285 GeneID:4318923 GeneID:4354193 GeneID:4321255 GeneID:4315909 |

4. Discussion

Among susceptible patients with impaired immune functions for a variety of reasons, such as solid organ transplant recipients and hematopoietic stem cell transplant recipients, *A. fumigatus*, *A. niger*, *A. terreus*, and *A. flavus* are the most common isolated species of IA, but the role of cryptic species should not be overlooked [29]. Species that appear to be cryptic might be neglected because of misidentification and therefore, it would be difficult to assess their epidemiological distribution pattern and antifungal susceptibility profile [5]. Antifungal susceptibility pattern, and rate and mechanism of resistance in ITC as a main agent in management and prophylaxis of aspergillosis against different species of *Aspergillus* have been analyzed in different countries [30–35]. In vitro activity of azoles determined by CLSI method against different opportunistic filamentous fungal pathogens including *Aspergillus* species from the Asia and Western Pacific Region, the data from the SENTRY Antifungal Surveillance Program (2011–2019) showed the ITC MIC$_{50/90}$ as 0.5/1 mg L$^{-1}$ for *A. fumigatus*, *A. flavus* and *A. terreus* and 1/2 mg L$^{-1}$ for *A. niger*. The overall frequency of NWT strains (MICs above the ECVs) of *A. fumigatus*, *A. flavus*, *A. terreus* and *A. niger* were 2.1%, 0, 0 and 2.2% for ITC, respectively [30]. They also reported an overall frequency of NWT strains of *A. fumigatus* as 0.9% for ITC. In addition, Europe displayed the highest NWT rates for ITC (1.7%) when compared to other regions (0.0–0.7%). During 2015 to 2019, Guegan et al. [32] analyzed a total of 929 *A. fumigatus* isolates from patients with cystic fibrosis forazole resistance. The rate of ITC resistant *A. fumigatus* isolates was 14.5% (95/656); they were recovered from 44/308 (14.3%) patients. The median MIC of ITC-resistant isolates was 14 mg/L$^{-1}$.

In a recently published study from Iran [33] MIC/MIC$_{50/90}$ values of ITC were reported for *Aspergillus* section *Fumigati* and *Nigri* as 4/≥8 µg mL$^{-1}$ (each) and 0.5/1 µg mL$^{-1}$ for *Aspergillus* section *Terrei*. They also no NWT strains reported in 233 *Aspergillus* isolates collected from 11 university hospitals in Iran. In addition, in two other studies from Iran, none of the *A. terreus* isolates [34] and different species of aspergillus including *A. tubingensis* and *A. niger* [35] had a MIC of ≥ ECV for ITC. In contrast to these studies, Nargesi et al. [3] reported a rate of 7.3% of NWT strains of *A. niger* against ITC. They also reported MIC$_{50/90}$ for *A. niger* and *A. tubingensis* against ITC as 0.25/0.5 and 0.5/16 µg mL$^{-1}$, respectively. In this current study, all tested *Aspergillus* species represented MICs under ECV against VOR, ISV and POS but MIC > 16 to ITC.
Despite all the measures that have been taken in the field of diagnosis of IA, the increasing prevalence of azole resistance has been found as a barrier in controlling this disease [36]. Small whole genome sequencing is a powerful technique which is used to capture variants throughout the entire genomic content of fungi and it provides a large amount of information in a short period of time compared to the approach target (such as Sanger sequencing) [37]. In azole-resistant strains of Aspergillus spp., the cyp51A and cyp51B genes may be exposed to polymorphism occurrence [13] The evidences gleaned from whole genome sequencing of Aspergillus paved the way for a clear description of how these genes operate without efforts to design multiple primers [14]. In regard to A. fumigatus, the majority of the existing data on tracking the effects of SNPs concentrate on defined sequences, such as the promoter region, TR34/L98H, of the cyp51A gene [38,39]. A. niger and A. tubingensis are two closely related species in the section Nigri [40]. The antifungal susceptibility profiles of these two species have already been reported in Iran [41] and in other countries [42]. A. fumigatus is more clear-cut in mechanisms of azole resistance than black aspergilli. Howard et al. [43] study on cyp51A gene mutations in multiple A. niger isolates showed contradictions in the concerned clades. They reported no mutation-related azole resistance in cyp51A gene in one clade, while six replacement mutations occurred in A. niger isolates of the other clade. Despite that, in our work, which overshadowed a larger scale of the genome, A. niger, with the exception of 15 silent mutations detected in the cyp51A and cyp51B genes, 3 and 2 replacement mutations were detected in cyp51A and cyp51B genes, respectively. Additionally, we detected 20 and 21 silent mutations in the cyp51B gene in A. tubingensis (isolates AT1 and AT2), respectively, despite the lack of mutations in the cyp51B gene (Table 2). A. welwitschiae is another member of the section Nigri [44]. Recent research showed that among the A. niger complex species, A. welwitschiae has a high isolation rate from clinical cases [39]. On the other hand, the antifungal susceptibility trend of A. welwitschiae has also been shown a higher MIC to VOR [45]. However, there is no exact information on how azole resistance has developed in A. welwitschiae. Based on the evidence from the whole genome sequencing of A. welwitschiae, substitutions at cyp51A and cyp51B, the two major target genes in A. welwitschiae isolate, as well as polymorphisms at multiple sites of target loci may explain the mechanism of ITC-resistance in our study. Zoran et al. [46] conducted a wide-range study on the azole resistance mechanism in Aspergillus terreus. Analysis of cyp51A gene, SNPs on 26 A. terreus isolates revealed that substitution at positions T650C, A649G, G1030A, and A956G resulted in changes in amino acid sequences. In another study performed by Rivero-Menendez et al. [47] on 12 triazole-resistant isolates of A. terreus, substitutions in D344N were perceived in two isolates and M217I in one isolate. In this study, between two isolates of A. terreus, only AR2 isolate showed a non-synonymous point mutation in the cyp51A and cyp51B genes separately (Table 2).

According to our findings, Aspergillus species isolated from different parts of the world may represent different pattern of resistant mechanisms including mutation in cyp51A and cyp51B genes which is clearly revealed by WGS. To show this, we made a literature review on the reported mutation in Aspergillus fumigatus revealed by Sanger sequencing of cyp51A and cyp51B genes (Table 3).

In this present study, we also evaluated non-cyp51 gene loci for mutations to study Aspergillus isolates. This is the first report from Iran. According to our findings, many mutations were observed in different isolates. The most isolates showed simultaneously mutations in cyp51 and non-cyp51 gene loci. However, one isolate of A. tubingensis (AT1) and A. terreus (AR2) did not show cyp51 mutations but AT2 and AR2 isolates showed mutations in AtrR, and Cdr1B and AtrR, respectively. A. fumigatus and AT1 (A. tubingensis) showed no mutations in cyp51B. However, they revealed mutations in non-cyp51 genes including AtrR and Hmg1, and Cdr1B, HapE and FfmA, respectively. It is worth to note that mutations in cyp51A gene associated with azole resistance are well known. However, different studies have been recently reported that some other gene loci including ATP-Binding Cassette Transporters (Cdr1B), AtrR, Hmg1, HapE and FfmA are required for wild-type azole
resistance in *A. fumigatus* as the most studied species in *Aspergillus* genus [17–21]. Likewise, it was suggested that the most common azole resistance allele currently described is a linked change corresponding to a change in the coding sequence of *cyp51A* and a duplication of a 34-bp region in the promoter leading to a tandem repeat [40, 48]. Paul et al. [18] also identified a positively acting transcription factor called *AtrR* that binds to the promoter of *cyp51A* as well as that of an important membrane transporter protein gene called *Cdr1B* (abcG1). These evidences have shown that non-*cyp51* gene loci such as *Cdr1B*, *AtrR*, *Hmg1*, *HapE* and *FfmA* are required for the synergistic increase in azole resistance and regulation of many different processes involved in drug resistance. Our finding on the homology modeling and mutation mappings of *CYP51A* of the selected *Aspergillus* species also suggest that the *cyp51A* may need other gene loci to cause azole resistance in *Aspergillus* species.

**Table 3.** An overview of the effects of point mutations on the emergence of azole resistance in *Aspergillus* spp. in the past decade.

| *Aspergillus* spp. | Target Genes | Point Mutations | Mutation Results | Authors (Year) [Reference] |
|-------------------|--------------|-----------------|------------------|-----------------------------|
| *A. fumigatus*    | Cyp51A       | TR34/L98H       | Observation of ITC-Resistant isolates + Observation of VOR-resistant isolates | Camps et al. (2012) [49] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H       | Observation of Triazole-Resistant isolates | Chowdhary et al. (2012) [50] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, TR46/Y121F/T289A | Observation of ITC-Resistant isolates Observation of VOR-resistant isolates | Astvad et al. (2014) [51] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H       | Observation of ITC-Resistant isolates | Ahmad et al. (2014) [52] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, TR46/Y121F/T289A | Observation of Triazole-Resistant isolates Observation of Triazole-Resistant isolates | van Ingen et al. (2015) [53] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, F46Y, D255E, and M172-G54E | Observation of Triazole-Resistant isolates Observation of Triazole-Resistant isolates -High MICs to both ITC and POS | Chowdhary et al. (2015) [54] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H       | Observation of ITC-Resistant isolates + Observation of VOR-resistant isolates | Nabili et al. (2016) [55] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, TR46/Y121F/T289A | Observation of ITC-Resistant isolates + Observation of VOR-resistant isolates | Wiederhood et al. (2016) [56] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, M220I, G448S, G54V | Observation of ITC-Resistant isolates | Mohammadi et al. (2016) [57] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H       | Observation of ITC-Resistant isolates | Hurst et al. (2017) [58] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, TR46/Y121F/T289A | Observation of ITC-Resistant isolates | Deng et al. (2017) [59] |
| *A. fumigatus*    | Cyp51A       | TR46/Y121F/T289A | Observation of ITC-Resistant isolates | Morio et al. (2018) [61] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H       | Observation of ITC-Resistant isolates | Paluch et al. (2019) [62] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, G448S, M220I | Observation of ITC-Resistant isolates | Tsuchido et al. (2019) [63] |
| *A. fumigatus*    | Cyp51A       | TR46/Y121F/T289A, G54V | Observation of ITC-Resistant isolates | Pontes et al. (2020) [64] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, G448S, M220I | Observation of VOR-Resistant isolates | Zhang et al. (2020) [65] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, G448S, M220I | Observation of VOR-Resistant isolates | Gonzalez-Jimenez et al. (2021) [58] |
| *A. fumigatus*    | Cyp51A       | TR34/L98H, G448S, M220I | Observation of VOR-Resistant isolates | Gonzalez-Jimenez et al. (2020) [16] |
| *A. fumigatus*    | Cyp51B       | G457S           | Observation of ITC-Resistant isolates + Observation of VOR-resistant isolates+ Observation of POS-resistant isolates+ Observation of ISV-resistant isolates | Gonzalez-Jimenez et al. (2020) [16] |
5. Conclusions

Our study indicates that cryptic species not only have a spread in clinical cases but also a high proportion of point mutations leading to azole resistance in their genomic composition. There is a possibility to say that this volume of mutations did not occur all at once in cryptic species, but the techniques of identifying these species have entered a new era and with the correct identification of species, researchers have been able to observe and confront the mutations that occur continuously. Further studies on the effect of mutations in \textit{cyp51} and non-\textit{cyp51} gene loci to cause clinical resistance in \textit{Aspergillus} isolates are warranted.

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