The A756T Mutation of the ERG11 Gene Associated With Resistance to Itraconazole in Candida Krusei Isolated From Mycotic Mastitis of Cows

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Candida krusei (C. krusei) has been recently recognized as an important pathogen involved in mycotic mastitis of cows. The phenotypic and molecular characteristics of 15 C. krusei clinical isolates collected from cows with clinical mastitis in three herds of Yinchuan, Ningxia, were identified by matrix-assisted laser desorption ionization–time of flight mass spectrometry. In addition to sequencing analysis, the ERG11 gene that encodes 14α-demethylases, the expression of the ERG11 gene, and efflux transporters ABC1 and ABC2 in itraconazole-susceptible (S), itraconazole-susceptible dose dependent (SDD), and itraconazole-resistant (R) C. krusei isolates was also quantified by a quantitative real-time reverse transcription polymerase chain reaction (qRT-PCR) assay. Sequencing analysis revealed three synonymous codon substitutions of the ERG11 gene including T939C, A756T, and T642C in these C. krusei clinical isolates. Among them, T642C and T939C mutations were detected in itraconazole-resistant and -susceptible C. krusei isolates, but the A756T substitution was found only in itraconazole-resistant isolates. Importantly, the expression of the ERG11 gene in itraconazole-resistant isolates was significantly higher compared with itraconazole-SDD and itraconazole-susceptible isolates (p = 0.052 and p = 0.012, respectively), as determined by the qRT-PCR assay. Interestingly, the expression of the ABC2 gene was also significantly higher in itraconazole-resistant isolates relative to the itraconazole-SDD and itraconazole-susceptible strains. Notably, the expression of ERG11 was positively associated with resistance to itraconazole (p = 0.4177 in SDD compared with S, p = 0.0107 in SDD with R, and p = 0.0035 in S with R, respectively). These data demonstrated that mutations of the ERG11 gene were involved in drug resistance in C. krusei. The A756T synonymous codon substitution of the ERG11 gene was correlated with an increased expression of drug-resistant genes including ERG11 and ABC2 in itraconazole-resistant C. krusei isolates examined in this study.

Keywords: Candida krusei, resistance, mycotic mastitis, cows, ERG11, ABC1, ABC2
INTRODUCTION

Cow mastitis has a major negative impact on dairy industries, causing significant economic losses to farmers. Pathogenically, a wide variety of microorganisms have been identified as causative agents of cow mastitis, mainly bacteria and fungi (1). In cases of fungal infection of the mammary gland, yeasts of Candida genus are the most reported fungal pathogens in cow mastitis (2).

Historically, the mycotic mastitis caused by fungi of the Candida genus was first described by Fleischer as early as 1930 (3). Although the infection of C. albicans was considered as the most common cause of mastitis, cases of cow mastitis caused by the infection of non-albicans Candida spp. (NAC), such as C. krusei, C. parapsilosis, C. glabrata, and C. tropicalis, have increased significantly during the last decade (4). Among these NACs, C. krusei ranked as the fifth most common cause of cow mycotic mastitis (5, 6). C. krusei was reported as the causative agent of bovine mastitis and bronchopneumonia in Canada, Mexico, Japan, the United Kingdom, Turkey, Poland, and Algeria (7, 8). Our previous investigation also suggested that C. krusei was one of the most important pathogens in mycotic mastitis in dairy farms of the Yinchuan region in Ningxia, China. This result was in accordance with the report of Erbaş et al. (9).

Candida krusei has been regarded as a multidrug-resistant fungal pathogen because of its intrinsic resistance to fluconazole (FLC) (10, 11), with more than 96% of human clinical and veterinary isolates being fluconazole-resistant (12). Azole is one of the most common antifungal drugs in agricultural practices (13), including itraconazole, ketoconazole, and tetracanazole. Although it has been reported that multiple mechanisms are involved in drug resistance in Candida spp., mechanisms involved in alterations of target enzymes and upregulation of multidrug resistance (MDR)-related proteins are the common mechanisms of Candida resistant to azoles. In this regard, 14α-lanosterol demethylase (14-DM) is the target enzyme of azoles, which is responsible for the production of an ergosterol precursor and is encoded by the ERG11 gene. In C. albicans and C. parapsilosis, the efflux pump genes CDR1, CDR2, and MDR1 are also associated with azole resistance (14). Nowadays, although transporter genes ABC1 and ABC2 were involved in drug resistance in C. krusei (15, 16), increased lines of evidence suggested that changes in the expression of activity of target enzyme and upregulation of MDR were the main mechanisms of drug resistance in C. krusei (11, 16). To date, the study on mechanisms of Candida in azole resistance has mainly focused on C. albicans, C. glabrata, and C. tropicalis, but studies on azole resistance mechanisms in C. krusei, especially the C. krusei isolates from cow mastitis resistant to itraconazole, are limited, and the involvement of ERG11, ABC1, and ABC2 genes in the drug resistance of C. krusei has not been determined (17).

In the present study, we evaluated the profile of the susceptibility of C. krusei to itraconazole and investigated the potential alterations of the ERG11 gene and the differential expression of ERG11, ABC1, and ABC2 genes of 15 clinical C. krusei isolates that were isolated from cow mastitis in Yinchuan, Ningxia, China.

MATERIALS AND METHODS

Isolation and Identification of Fungal Pathogens

This study was submitted to and approved by the Ethic Committee of Animal Study in Ningxia University. A total of 465 quarter-milk samples were collected from the cows with clinical or subclinical mastitis, which originated from three herds in Yinchuan, Ningxia, China. Clinical or subclinical mastitis was defined by swelling, reduced milk flow, and abnormal milk appearance (watery to viscous with clots varying from gray-white to yellowish). Additionally, other signs of infection such as fever, inappetence, ataxia, and depression were also considered. These cows have been treated with antibiotics before sample collection. The isolates of C. krusei were identified by using Candida chromogenic medium (CHROMagar, France) (18) and matrix-assisted laser desorption ionization–time of flight mass spectrometry (MALDI-TOF MS) with score values of >2.000 (VITEK® MS, BioMerieux, France) (19) and stored in liquid nitrogen.

Antimicrobial Susceptibility Tests

The susceptibility assay was conducted using the Clinical and Laboratory Standard Institute (CLSI) broth microdilution (BMD) method. The CLSI BMD method was performed in a 96-well polystyrene microtiter plate in accordance with CLSI M27-A3 and M27-S4 guidelines (20, 21). The ranges of concentrations of tested drugs were as follows: 5-flucytosine (0.03–64 µg/ml), amphotericin B (0.008–16 µg/ml), fluconazole (0.03–64 µg/ml), itraconazole (0.03–64 µg/ml), and ketoconazole (0.03–64 µg/ml) (15, 22). The antifungal drugs were all purchased from Meilun Biotechnologies (Dalian, China). The C. krusei NCCLS reference strain ATCC 6258 served as quality control to ensure the test (3, 23); 1 × 10^5 CFU/ml working suspension of the C. krusei isolates was added into the each well. Results were recorded as resistant, susceptible dose dependent, and sensitive as shown in Table 1.

PCR Amplification and Sequencing Alignment of the ERG11 Gene

Candida krusei isolates were subcultured twice on Sabouraud agar at 37°C for 18–24 h to revive and ensure the purity of cultures. A single colony was then transferred to 20 ml of liquid YPD (yeast extract 1%, dextrose 2%, and peptone 2%) broth and cultured at 35°C in a shaking incubator (120 rpm) exponential growth phase. The bacteria cells were collected by centrifugation at 3,000 rpm for 20 min, and the bacteria pellet was used for total genomic DNA preparation using a DN515-A new Plant Genomic DNA Rapid Extraction Kit (Aidlab Biotechnologies, Beijing, China) according to the manufacturer’s instruction. The isolated DNA was used as a template for amplification of the ERG11 gene. The primer set of ERG11 was designed by Primer 5.0 and synthesized at Sangon Biotech (Shanghai, China), based on the available sequence information of the C. krusei ERG11 gene (Gene accession number DQ903905) at the National Center for Biotechnology Information (NCBI) (Table 2). The PCR amplification was conducted in 25 µl volume containing
| Name          | Susceptibility profile |
|--------------|-----------------------|
| ATCC 6258    | ITR: 0.06/S, AMB: 0.5/S, FLC: 8/S, 5-FC: 4/S, KET: 0.125/S |
| CK1          | ITR: 4/R, AMB: 2/SDD, FLC: 64/R, 5-FC: 32/R, KET: 1/R |
| CK2          | ITR: 8/R, AMB: 2/SDD, FLC: 64/R, 5-FC: 32/R, KET: 1/R |
| CK3          | ITR: 0.25/SDD, AMB: 2/SDD, FLC: 16/SDD, 5-FC: 32/R, KET: 1/R |
| CK4          | ITR: 0.25/SDD, AMB: 2/SDD, FLC: 64/R, 5-FC: 32/R, KET: 0.25/SDD |
| CK5          | ITR: 8/R, AMB: 4/R, FLC: 64/R, 5-FC: 32/R, KET: 1/R |
| CK6          | ITR: 0.5/SDD, AMB: 2/SDD, FLC: 64/R, 5-FC: 32/R, KET: 1/R |
| CK7          | ITR: 0.06/S, AMB: 2/SDD, FLC: 16/SDD, 5-FC: 32/R, KET: 0.25/SDD |
| CK8          | ITR: 0.25/SDD, AMB: 2/SDD, FLC: 64/R, 5-FC: 32/R, KET: 1/R |
| CK9          | ITR: 4/R, AMB: 2/SDD, FLC: 64/R, 5-FC: 32/R, KET: 1/R |
| CK10         | ITR: 0.25/SDD, AMB: 2/SDD, FLC: 32/SDD, 5-FC: 32/R, KET: 1/R |
| CK11         | ITR: 0.25/SDD, AMB: 2/SDD, FLC: 64/R, 5-FC: 16/SDD, KET: 0.5/SDD |
| CK12         | ITR: 4/R, AMB: 4/R, FLC: 64/R, 5-FC: 32/R, KET: 1/R |
| CK13         | ITR: 0.5/SDD, AMB: 2/SDD, FLC: 64/R, 5-FC: 32/R, KET: 1/R |
| CK14         | ITR: 0.06/S, AMB: 2/SDD, FLC: 16/SDD, 5-FC: 32/R, KET: 0.5/SDD |
| CK15         | ITR: 0.25/SDD, AMB: 2/SDD, FLC: 64/R, 5-FC: 16/SDD, KET: 1/R |

CK, Candida krusei; S, susceptible; SDD, susceptible dose dependent; R, resistant; 5-FC, 5-flucytosine, MIC break point: S, ≤0.125 µg/ml; R, ≥1 µg/ml; AMB, amphotericin B, MIC break point: S, ≤1 µg/ml; R, ≥2 µg/ml; FLC, fluconazole, MIC break point: S, ≤8 µg/ml; R, ≥32 µg/ml; ITR, itraconazole, MIC break point: S, ≤0.125 µg/ml; R, ≥1 µg/ml; KET, ketoconazole, MIC break point: S, ≤0.125 µg/ml; R, ≥2 µg/ml.

1 µl of genomic DNA (200 ng/µl), 0.5 µl of specific forward and reverse primers (50 µmol/L), and 12.5 µl of 2× Phanta Max Master Mix (Vazyme Biotech, Nanjing, China). The PCR parameters were set as denaturation for 3 min at 95°C, followed by 35 cycles of 95°C for 15 s, 60°C for 15 s, and 72°C for 1 min, and a final step of elongation (72°C for 5 min). The resultant PCR product was cleaned by gel purification in 1.5% agarose prior to being cloned into the pTOPO-TA Vector using a CV16-Zero Background pTOPO-Blunt Cloning Kit with Blue/white selection (Aidlab Biotechnologies, Beijing, China). The white colonies were analyzed for clones containing the DNA fragment of gene of interest. Eight to fifteen plasmids from clones generated from an identical PCR product were further sequenced for the ERG11 gene (Sangon Biotech, Shanghai, China). The sequences were aligned with the online published sequence of the ERG11 gene (Gene Accession Number DQ903905) to determine gene mutation (17).

Quantitative Real-Time PCR Analysis
For quantitative real-time PCR (qRT-PCR) analysis, total RNA was extracted from C. krusei cultures with RNAiso Reagent (TaKaRa, Dalian, China) and reverse transcribed to cDNA with HiScript III RT SuperMix for qPCR (+gDNA wiper) (Vazyme Biotech, Nanjing, China) according to the manufacturer’s instruction. For the ERG11 target gene and GAPDH reference gene, primer pairs were designed with the Primer 5.0 program and synthesized by Sangon, Shanghai, China (Table 2). qRT-PCR was conducted with a 20-µl volume containing the following reagents: 10 µl of 2× ChamQ Universal SYBR qPCR Master Mix (Vazyme), 1 µl of total RNA sample, 0.5 µl of each primer pair at a concentration of 10 µmol/L, and 8 µl of distilled water. Each reaction was run in triplicate. Samples were subjected to an initial step at 95°C for 3 min, followed by 40 cycles, each of which consisted of 10 s at 95°C and 30 s at 60°C. Melting curves were recorded every 5 s. The fluorescence data were collected and analyzed with the QuantStudio Design Analysis Software 1.3.1. A 2−ΔΔCt algorithm was employed to analyze the relative expression levels of drug-resistant genes at resistant, susceptible dose-dependent, and sensitive strains.

Statistical Analysis
Statistical analysis was performed with the GraphPad Prism program (GraphPad 8.0.1 Software Inc., San Diego, CA). The two-tailed Student’s t-test was used to analyze significant differences between gene expression displayed by the distinct C. krusei strains; p < 0.05 was considered statistically significant.

RESULTS
Resistance of C. krusei Isolates to Antifungal Agents
A total of 15 C. krusei isolates (designated as CK1–CK15) were isolated from clinical samples from April 2018 to October 2019 in three herds at Yinchuan, Ningxia, China. Drug sensitivity testing was performed according to the broth microdilution method M27-A2 (NCCLS 2002), and the result showed that among 15 C. krusei isolates, 73.4, 73.4, and 66.7% were resistant to fluconazole (FLC), ketoconazole (KET), and 5-flucytosine (5-FC), respectively. However, 20 and 33.3% of the isolates were susceptible to amphotericin B (AMB) and itraconazole (ITR), respectively. Interestingly, isolates CK2, CK5, CK6, CK8, CK9, CK12, and CK13 showed resistance to both 5-FC and FLC.
missense mutation might not be associated with drug resistance. All sequenced isolates, except the reference strain ATCC 6258, were found to harbor a missense mutation (C44T) in the ERG11 gene. This alteration resulted in an amino acid change from alanine to valine. However, such a missense mutation was also found in the reference strain ATCC 6258, which indicated that the C44T missense mutation might not be associated with drug resistance to azoles in C. krusei.

**Mutational Analysis in ERG11 of C. krusei Isolates**

The ERG11 gene fragment was amplified from ATCC 6258 and all 15 C. krusei isolates. The PCR product of the open frame of the ERG11 gene was 1,587 bp, which encodes 529 amino acids. Sequencing analysis identified four different mutations, three synonymous mutations (C642T, A756T, and T939C), and one missense mutation (C44T) in these 15 C. krusei isolates (Table 4). Synonymous mutations C642T and T939C were presented in 44 bp (C → T) of the ERG11 gene (Table 4), which resulted in an amino acid alteration from alanine to valine. However, such a missense mutation was also found in the reference strain ATCC 6258 and all C. krusei strains, which indicated that the C44T missense mutation might not be associated with drug resistance to azoles in C. krusei.

**Increased ERG11 Gene Transcript in Itraconazole-Resistant C. krusei**

In order to examine whether an alteration of ERG11 gene expression was correlated with the drug resistance of C. krusei clinical isolates, the transcript of the ERG11 gene was accessed by a qRT-PCR assay. In comparison with the reference strain ATCC 6258, the relative ERG11 gene expression of field isolates in five itraconazole-resistant isolates was significantly upregulated (p < 0.01), while only two itraconazole-susceptible dose-dependent isolates showed a significantly upregulated ERG11 gene expression (p < 0.01, Figure 1A). The result showed that the transcript of the ERG11 gene in itraconazole-resistant isolates was significantly more abundant than itraconazole-susceptible strains (p = 0.0012, Figure 1B) and itraconazole-susceptible dose-dependent (SDD) strains (p = 0.0052, Figure 1B). However, there was no significant difference between itraconazole-susceptible dose-dependent isolates and itraconazole-susceptible isolates (p = 0.2562, Figure 1B).

**Increased ABC2 Gene Transcript in Itraconazole-Resistant C. krusei**

ABC transporters are involved in drug resistance; next, we therefore sought to examine the alteration of ABC transporters in C. krusei isolates. Interestingly, unlike the ERG11 gene, none of field isolates showed an upregulated ABC1 gene expression as compared with that of the reference strain.
TABLE 4 | ERG11 gene point mutations in C. krusei clinical isolates.

| Information of strains | ERG11 gene mutation sites |
|------------------------|---------------------------|
| Name                   | ITR susceptibility category | 44 | 642 | 756 | 939 |
| DQ903905               | S                         | T  | T  | A  | T  |
| ATCC 6258              | R                         | C  | C  | —  | —  |
| CK1                    | R                         | C  | —  | T  | C  |
| CK2                    | —                         | —  | —  | T  | C  |
| CK3                    | SDD                       | C  | —  | —  | —  |
| CK4                    | SDD                       | —  | —  | —  | —  |
| CK5                    | R                         | —  | C  | T  | C  |
| CK6                    | SDD                       | C  | —  | —  | —  |
| CK7                    | S                         | C  | —  | —  | —  |
| CK8                    | SDD                       | —  | —  | —  | —  |
| CK9                    | R                         | —  | —  | T  | —  |
| CK10                   | SDD                       | C  | C  | —  | —  |
| CK11                   | SDD                       | —  | —  | —  | —  |
| CK12                   | R                         | —  | —  | T  | C  |
| CK13                   | SDD                       | C  | —  | —  | —  |
| CK14                   | S                         | —  | C  | —  | C  |
| CK15                   | SDD                       | C  | —  | —  | —  |

The numbering of the nucleotides shown in this table starts with 1 for the A of the ATG start codon. DQ903905, GenBank Accession no. of C. krusei whose ERG11 sequence published online is used to align with C. krusei clinical isolates in this study.

ATCC 6258 (Figure 2A). The results of qRT-PCR for ABC1 genes showed that the expression of ABC1 gene mRNA was not significantly different between itraconazole-resistant isolates, itraconazole-susceptible dose-dependent isolates, and itraconazole-susceptible strains (p = 0.3844, p = 0.9997, and p = 0.2996, respectively, Figure 2B). Similar to that seen in the ERG11 gene, the relative ABC2 gene expression was extremely significantly upregulated in four itraconazole-resistant isolates (p < 0.01) and significant in one itraconazole-resistant isolate (p < 0.05), as compared with the reference strain ATCC 6258, whereas only two itraconazole-susceptible dose-dependent isolates showed an extremely significant upregulation of ABC2 gene expression compared to the reference strain ATCC 6258 (p < 0.01, Figure 3A). Intriguingly, the transcript
of the ABC2 gene in itraconazole-resistant isolates was more abundant relative to itraconazole-susceptible strains ($p = 0.0035$) and itraconazole-susceptible dose-dependent strains ($p = 0.0107$, Figure 3B). No significant difference was found in the expression of the ABC2 gene was determined between the itraconazole-susceptible dose-dependent group and itraconazole-susceptible *C. krusei* isolates ($p = 0.4177$, Figure 3B).

**DISCUSSION**

Our previous study demonstrated that *C. krusei* was a predominant pathogen isolated from mycotic mastitis of cows in Yinchuan, Ningxia, China (12), suggesting that it may be an important fungal pathogen of mycotic mastitis of cows in this area. The crucial roles of drug-resistant genes ERG11, ABC1, and ABC2 in FLC-resistant clinical isolates of *C. krusei*
from human have been well-established (15, 24). However, the pathogenic molecular mechanism of *C. krusei* isolated from cow mastitis remains unclear. In the present report, we evaluated the susceptibility profiles and mutations in the *ERG11* gene in 15 clinical *C. krusei* isolates. The expression of drug-resistant genes *ERG11*, *ABC1*, and *ABC2* between isolates susceptible, susceptible dose-dependent, and resistant to ITR was also analyzed. We identified three synonymous mutations and one missense mutation in the *ERG11* gene in these clinical *C. krusei* isolates, as previously described in human *C. krusei*. Furthermore, the A756T was only presented in ITR-resistant strains, suggesting that it might be correlated with drug resistance in *C. krusei*, while mutations T642C and T939C were presented in all these 15 *C. krusei* isolates. The expression of drug-resistant *ERG11* and *ABC2* was also significantly higher in ITR-resistant *C. krusei* isolates compared to ITR-susceptible and susceptible dose-dependent isolates, suggesting a correlation of mutations of the *ERG11* gene with the resistance to antifungal agents in *C. krusei*.

In the present study, based on the CLSI BMD method's susceptibility to ITR, the 15 *C. krusei* clinical isolates from cows with clinical mastitis could be divided into three groups: ITR-susceptible (2 isolates), ITR-susceptible dose-dependent (8 isolates), and ITR-resistant (5 isolates). The antifungal testing showed that 13.3, 53.3, and 33.3% were susceptible, susceptible dose-dependent, and resistant to itraconazole among these *C. krusei* clinical isolates, respectively. Notably, 13 of the 15 *C. krusei* isolates (73.4%) were also resistant to FLC and KET, 10 of the 15 *C. krusei* isolates (66.7%) were resistant to flucytosine, and 3 of the 15 isolates (20%) were resistant to amphotericin B. This finding was consistent with a study by Namvar et al. (25), but was different from reports by others (8, 9, 26). Of interest, the rate of resistance to antifungal agents in these *C. krusei* isolates was lower than our previous findings (12), which might be attributed to the reduction in the use of antifungal drugs during breeding. Consistent with our previous studies, *C. krusei* isolates were double-resistant and multidrug-resistant to antifungal drugs. It is strongly recommended that ketoconazole and other azole antifungal agents should not be used in the treatment of *C. krusei* infection in dairy cows in Ningxia, China, owing to high drug resistance.

It is worth noting that CK2, CK5, and CK12 showed resistance to amphotericin B, but such cases are rare (27). However, several lines of studies evidenced an increased minimum inhibitory concentration (MIC) of amphotericin B in *C. krusei* isolates. In *Candida* spp., resistance to amphotericin B was found to be associated with a decreased ergosterol content of cell membrane (28–30). In addition, an inactivation of *ERG3* could substitute 14α-methylcholesterol with ergosterol, thus reducing ergosterol levels, which, in turn, resulted in deficient ergosterol to counter the function of amphotericin B (31). Moreover, several other mutations in the ergosterol biosynthetic genes such as *ERG2*, *ERG5*, *ERG6*, and *ERG24* can also result in *C. albicans* and *C. glabrata* resistant to amphotericin B (32–34). However, such mutations have not been reported in *C. krusei* and further studies are needed in future studies.

A number of studies have focused on *ERG11* gene mutation in *Candida* species (35–37). In order to fully understand *ERG11* gene mutation of *C. krusei*, the whole open reading frame of the *ERG11* gene was amplified for sequencing analysis, and three synonymous mutations and one missense mutation were identified in this study. C44T has one missense mutation and was found in all 15 isolates and the reference strain, suggesting that it had no impact on the itraconazole resistance of *C. krusei*. Molecularly, the C447T mutation resulted in the alteration of alanine to valine in the 15th amino acid of 14α-lanosterol demethylase (14-DM); this mutation might occur outside the active site of the *ERG11* gene, which might not affect the mutual interaction of azole and 14-DM, or a single missense mutation might not be sufficient to change the affinity of the 14-DM to azole (15). Moreover, synonymous mutations (C642T, A756T, and T939C) in the 15 isolates were consistent with a previous report (24). Synonymous mutations can affect transcription, splicing, mRNA transport, and translation, any of which could change phenotype, rendering the synonymous mutation non-silent (38). He et al. (15) also reported that C642T, T1389C, and G1536C mutations occurred in all the experimental strains. In addition, the T1389C mutation was also reported by Ricardo et al. (16). Tavakoli et al. (35) revealed a heterozygous polymorphism at position T939C in the *ERG11* coding region and speculated that this polymorphism might play a key role in the transcriptional regulation of genes and be involved in the processes of ergosterol biosynthesis. Of note, several previously reported mutations, including the T418C missense mutation (16), and C51T, T1389C, and G1536C synonymous mutations (15), have not been found in this study; thus, limited clinical isolates (15) were analyzed. Mechanistically, previous studies on *C. albicans* and *C. tropicalis* have demonstrated that the missense mutation was associated with resistance to azole, which was partly through changing the conformation of the target enzyme 14-DM, which, in turn, decreased its drug affinity and influenced the enzyme's function in ergosterol biosynthesis (39, 40). The resistance mechanism of these resistant strains may be due to one or multiple mutations in these genes, which needs further investigation.

In addition, the expression of *ABC2* and *ERG11* genes was significantly upregulated in *C. krusei* veterinary clinical isolates resistant to itraconazole. Previous studies demonstrated that resistance to azole was also due to the increased expression of *ERG11*. This results in insufficientazole activity owing to the overproduction of the target enzyme (41). Although *ERG11* overexpression has been reported in *C. krusei*, the mechanism behind the overexpression remains unclear (15, 35). Another mechanism of resistance to azole is via the decreased intracellular accumulation of azole. This can be due to efflux pump activity or changes in the cell membrane. In this regard, drug efflux pumps belong to either the ATP-binding cassette (ABC) family of transporters or the major facilitator super family (MFS) class. These proteins can pump out fungicidal compounds across the cell membrane, and their overexpression results in multidrug resistance phenotype in pathogenic fungus (17). In contrast to members of the MFS class that are actuated by electrochemical proton-motive force, ABC family members depend on the hydrolysis of ATP for energy (42, 43). Indeed, along with the decrease of susceptibility to itraconazole, the *ABC2* gene mRNA
expression in the isolates appeared to increase. Although it was considered that Abc1p played an important role in the innate FLC resistance of \textit{C. krusei} (16, 17), the ABC2 gene could be activated slower than the \textit{ABC1} gene in the presence of voriconazole (16). Given voriconazole tolerance, the Abc1p efflux pump is supposed to be more efficient in expelling drug and plays a late role in the development of resistance, and the accumulation of itraconazole in drug-susceptible \textit{C. krusei} was higher than that of resistant strains (44). Interestingly, Venkateswarlu et al. found that two isolates highly resistant to fluconazole showed a different sensitivity to itraconazole, suggesting that itraconazole and fluconazole in \textit{C. krusei} may have different resistance mechanisms (44). The authors stated that \textit{C. krusei} resistant to itraconazole was due to decreased drug accumulation in cells and speculated that there might exist more efflux pumps contributing to itraconazole resistance of \textit{C. krusei}, which could be well-explained in this study. The overexpression of both \textit{ERG11} and \textit{ABC2} has been reported to be involved in itraconazole resistance in \textit{C. krusei} (15, 35). However, an unusual transient or stable resistance of \textit{C. krusei} to voriconazole has also emerged. Overexpression of \textit{ABC2} and \textit{ERG11}, as observed in itraconazole resistance, imparts a transient resistance to voriconazole, while a more stable resistance was observed due to the overexpression of \textit{ABC1} and point mutation in \textit{ERG11} (16). Taking this into account, we speculate that the resistance mechanisms of itraconazole and voriconazole in \textit{C. krusei} clinical isolates may be different. Moreover, other genes encoding ATP-dependent efflux transporters may occur in \textit{C. krusei}, such as a CgSNQ2 homologous gene that was verified as anazole-associated resistance gene in \textit{C. glabrata} (45). Although the \textit{C. krusei} genome has been sequenced, it is not completely annotated yet; thus, other transporter genes were not assessed. Our mRNA expression data showed that Abc2p may play a more important role in itraconazole resistance of \textit{C. krusei}, instead of Abc1p.

This study has enriched our knowledge in the veterinary clinical \textit{C. krusei} resistance gene expression and mutation data by comparing the difference between the veterinary clinical and the human clinical \textit{C. krusei}, and further deepened our understanding of the resistance mechanism of \textit{C. krusei} in veterinary clinics. The limitation of this study is that the sample size was small and no drug susceptibility test and resistance mechanism research related to echinocandin has been included, which require further investigations.

In conclusion, in this study, we found that \textit{C. krusei} veterinary clinical isolates exhibited a different susceptibility to antifungal agents. Mechanistically, the A756T mutation in the \textit{ERG11} gene resulted in an upregulation of drug-resistant genes \textit{ERG11} and \textit{ABC2}, substantially enhancing the resistance to itraconazole of \textit{C. krusei}. Although we have identified four point mutations in the \textit{ERG11} gene associated with itraconazole resistance and have already described their role on the itraconazole resistance of \textit{C. krusei}, it is necessary to confirm the effect of these mutations by site-directed mutagenesis of the \textit{C. krusei} strain in the future. Nevertheless, this study may thus provide an insight into the mechanism of the resistance of \textit{C. krusei} to antifungal agents, which warrants for further investigation.

**DATA AVAILABILITY STATEMENT**

The datasets generated for this study are available on request to the corresponding author.

**ETHICS STATEMENT**

The animal study was reviewed and approved by Ethics Committee of Animal Research of Ningxia University.

**AUTHOR CONTRIBUTIONS**

JD, YW, and XZ conceived and designed the experiments. JD and JF analyzed the data and drafted the manuscript. JD, WM, JF, and XL performed experiments and acquired data. XL and XZ interpreted data and critically revised the manuscript. All authors read and approved the final version of the manuscript.

**FUNDING**

This study was supported by a grant from the National Natural Science Foundation of China (No. 32060816), a grant from the Key Research and Development Plan Project of Ningxia Hui Autonomous Region (No. 2017BN04), and a grant from Natural Science Foundation of Ningxia (2019AAC03006). These funds play no role in the design of the study; collection, analysis, and interpretation of data; and the writing of the manuscript.

**ACKNOWLEDGMENTS**

The authors thank all the study participants, farm owners, veterinarians, and personnel of the Center of Disease Control and Veterinary Institute in Ningxia Hui Autonomous Region of China, who helped in the realization of this study.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.