Transcriptome sequencing revealed the inhibitory mechanism of ketoconazole on clinical Microsporum canis

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ABSTRACT

Background: Microsporum canis is a zoonotic disease that can cause dermatophytosis in animals and humans.

Objectives: In clinical practice, ketoconazole (KTZ) and other imidazole drugs are commonly used to treat M. canis infection, but its molecular mechanism is not completely understood. The antifungal mechanism of KTZ needs to be studied in detail.

Methods: In this study, one strain of fungi was isolated from a canine suffering with clinical dermatosis and confirmed as M. canis by morphological observation and sequencing analysis. The clinically isolated M. canis was treated with KTZ and transcriptome sequencing was performed to identify differentially expressed genes in M. canis exposed to KTZ compared with those unexposed thereto.

Results: At half-inhibitory concentration (½MIC), compared with the control group, 453 genes were significantly up-regulated and 326 genes were significantly down-regulated (p < 0.05). Quantitative reverse transcription polymerase chain reaction analysis verified the transcriptome results of RNA sequencing. Gene ontology enrichment analysis and Kyoto Encyclopedia of Genes and Genomes enrichment analysis revealed that the 3 pathways of RNA polymerase, steroid biosynthesis, and ribosome biogenesis in eukaryotes are closely related to the antifungal mechanism of KTZ.

Conclusions: The results indicated that KTZ may change cell membrane permeability, destroy the cell wall, and inhibit mitosis and transcriptional regulation through CYP51, SQL, ERG6, ATM, ABCB1, SC, KER33, RPA1, and RNP genes in the 3 pathways. This study provides a new theoretical basis for the effective control of M. canis infection and the effect of KTZ on fungi.

Keywords: Microsporum canis; transcriptome; ketoconazole

INTRODUCTION

Microsporum canis is a ubiquitous protozoan pathogenic fungus [1,2]. M. canis mainly infects dogs, cats, and other animals, and then the animal infects humans [3]. Children and
immunocompromised individuals are more readily infected [4]. The main symptoms are hair loss, tinea capitis, tinea pedis, and onychomycosis [1,5]. Currently, various oral and topical antifungal drugs such as griseofulvin, terbinafine, itraconazole (IT), fluconazole (FLZ), and imidazole drugs are commonly used to control M. canis infection [6], among which ketoconazole (KTZ), as an imidazole drug, is the most frequently used [7]. Imidazoles are synthetic antifungal drugs that can selectively inhibit fungal cytochrome P-450-dependent 14-α-demethylase, change the permeability of cell membranes, causing the loss of important intracellular substances, leading to fungal death [8]. Imidazole exhibits selective toxicity to fungi and this is mainly related to the inhibition of ergosterol biosynthesis and interference with other membrane lipids [9,10]. Although the antifungal mechanism of KTZ is similar to other imidazole drugs, its potential antifungal mechanism warrants further elucidation.

Through transcriptome sequencing technology, it is possible to obtain almost all transcript sequence information of a species in a certain state [11,12]. Transcriptome research allows the study of gene function and gene structure at a holistic level, revealing specific biological processes, and has been widely used in basic research, clinical diagnosis, and drug research and development [11-13]. In this study, we utilised transcriptome sequencing to explore differentially expressed genes (DEGs) and the pathways regulated by DEGs of M. canis after KTZ exposure, which clarified the underlying molecular mechanisms, signalling pathways of inhibition of KTZ on M. canis. Our study provides a new theoretical basis for the effective control of M. canis infection and the mechanism of KTZ inhibition on fungi.

MATERIALS AND METHODS

Cultivation and identification of M. canis

M. canis samples were taken from the skin of dogs suffering from skin diseases from an animal hospital in Shenyang, China. Subsequently, clinical samples were cultured using Sabouraud’s Dextrose Agar medium (Coolaber, China), and colonies similar to the morphology of M. canis were isolated and purified by using potato dextrose agar (PDA) medium (Solarbio, China). Subsequently, Tryptone Soy Broth (Solarbio) was cultured in a fungus incubator at 28°C for 96 h, and the concentration of the fungal solution was adjusted to 2 × 10⁴ CFU/mL. KTZ was purchased from Solarbio Ltd. (Solarbio) and dissolved in dimethyl sulfoxide (DMSO). The experiment was divided into 2 groups: the test group (T4,T5, and T6) treated with KTZ at a concentration of ½MIC for 6 h, and the control group (T1,T2, and T3) treated with DMSO at the same concentration. Tests on specimens in each group were repeated 3 times.

Identification of M. canis

The purified and cultured fungi were subjected to Diff-Quik® staining, morphological observation, and 18s ribosomal DNA (rDNA) sequence analysis. The DNA of M. canis was extracted using the Ezup column fungal genomic DNA extraction kit (Sangon Biotech Co., Ltd., China). Polymerase chain reaction (PCR) was carried out to amplify the DNA of M. canis. The primer sequences used for PCR were (NS1: f:5′-GTAGTCATATGCTTGTCTC-3′ and NS6: 5′-GCACTACACAGACCTGATTTGCGCTC-3′). PCR was performed under following conditions: a hot start at 95°C for 4 min, then 30 cycles of 94°C (45 sec), 55°C (45 sec), and 72°C (60 sec), followed by extension at 72°C for 10 min. The resulting PCR amplification products were sequenced by Sangon Biotech Co., Ltd. BLAST (http://www.ncbi.nlm.nih.gov) was used to
identify sequences similar to the *M. canis* gene. All similar sequences were found from NCBI and a phylogenetic tree built after DNASTAR's alignment.

**The minimum inhibitory concentration of KTZ against *M. canis***

*M. canis* and RPMI1640 medium were thoroughly mixed, centrifuged, and the fungal suspension obtained; the concentration of the RPMI1640 fungal suspension was adjusted to \(4 \times 10^4\) CFU/mL. According to the CLSI M27-A standard, a drug susceptibility test was carried out on *M. canis* by micro-broth dilution method. We inoculated the fungal solution into a 96-well plate, dissolved KTZ with DMSO and added it to the fungal solution so that the final concentration of KTZ therein was 0 \(\mu\)g/mL, 0.195 \(\mu\)g/mL, 0.39 \(\mu\)g/mL, 0.78 \(\mu\)g/mL, 1.56 \(\mu\)g/mL, 3.13 \(\mu\)g/mL, 6.25 \(\mu\)g/mL, 12.5 \(\mu\)g/mL, and 25 \(\mu\)g/mL. The same concentration of DMSO without KTZ was used as a negative control. The 96-well plate was placed in an incubator and held at 28°C for 7 days whereafter the optical density value was measured.

**Library preparation for RNA and transcriptome sequencing**

Total RNA was isolated from samples using the TRUEscript RT MasterMix (XinBaiji Biotech, China) according to manufacturer’s instructions. According to the manufacturer’s instructions, NanoDrop 2000 (Thermo Fisher Scientific, USA) was used to measure the RNA concentration and purity, and the Agilent Bioanalyzer 2100 system (Agilent Technologies, USA) was employed to identify RNA integrity.

After the total RNA passed quality inspection, the magnetic beads connected to Oligo (dT) were used to enrich the eukaryotic messenger RNA (mRNA). The extracted mRNA was randomly interrupted into short fragments by RNA fragmentation buffer. Using fragmented mRNA as a template, a single-stranded complementary DNA (cDNA) was synthesised using random hexamers with 6 bases. Then buffer, dNTPs, RNaseH, and DNA polymerase I (Pol I) were added for double-strand cDNA synthesis. AMPure XP beads were used to purify double-stranded products, and the activities of T4 DNA polymerase and Klenow DNA polymerase were used to repair the sticky ends of DNA to form blunt ends. AMPureXP beads were used to perform fragment selection. Finally, PCR amplification was performed to obtain the final sequencing library. After the library was qualified, the Illumina Hiseq4000 was used for sequencing.

**Data analysis**

Cutadapt and internal Perl scripts were utilised to delete reads that contain adapter contamination and low-quality or uncertain bases. FastQC ([www.bioinformatics.babraham.ac.uk/projects/fastqc/](https://www.bioinformatics.babraham.ac.uk/projects/fastqc/)) was used to identify the quality of the sequence, including determining the quality scores Q20 and Q30 and the GC content of the clean data. All downstream analyses used high-quality clean data. Then we compared the high-quality sequence obtained after quality control with the designated reference genome.

**Differential expression analysis**

Differential expression analyses of 2 groups were performed using the DESeq2. The resulting \(p\) values were adjusted using the Benjamini and Hochberg’s approach for controlling the false discovery rate. Genes with an adjusted \(p\) value of less than 0.01 found by DESeq2 were assigned as differentially expressed.

**Gene ontology (GO) and KEGG pathway enrichment analyses**

GO and KEGG databases were used for enrichment analysis of DEGs to ascertain the functions of DEGs. The Goseq R software package was used for the GO enrichment analysis
of DEGs. KOBAS software was employed to test the statistical enrichment of DEGs in the KEGG pathway.

**Verification of DEGs**

Total RNA was isolated from samples using TRUEscript RT MasterMix (XINBAJII Biotech) according to manufacturer’s instructions. The A260/A280 ratio of RNA was detected using an ultra-micro-nucleic acid protein analyser (Analytik Jena AG, Germany). When the A260/A280 ratio of RNA ranged from 1.8 to 2.0, it was used in subsequent experiments.

According to the instructions provided by the manufacturer, the first strand of cDNA was synthesised using TransScript First-Strand cDNA Synthesis kit (AiDLAB Biotech, China). Thereafter, according to the manufacturer’s requirements, quantitative reverse transcription PCR (qRT-PCR) was conducted on an ABI 7500 real-time PCR system using SYBR Green QPCR Mix (DF Biotech, China) under the following conditions: thermal cycling denaturation at 95°C for 3 min, followed by 40 cycles at 95°C for 10 sec, annealing at 60°C for 30 sec, and then extension. PheRS can be used as an internal reference gene. Finally, the relative change in mRNA was calculated using the $2^{-\Delta\Delta Ct}$ method. **Table 1** displays the primers used in the experiment. The primers were synthesised by Sangon Biotech Co., Ltd.

**Statistical analysis**

All statistical tests were conducted using SPSS 19.0 software (IBM, USA) and the results were expressed as mean ± standard deviation ($X \pm SD$). To reveal the difference between 2 groups, 2-tailed Student’s $t$-tests were used and significant differences among multiple groups were evaluated by using 1-way analysis of variance and least significant difference methods. Differences of $p < 0.05$ were considered significant.

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**Table 1.** Primer sequences of messenger RNA used for quantitative real-time polymerase chain reaction

| Gene name | Primer sequence (5'-3') | Annealing temperature (°C) |
|-----------|-------------------------|----------------------------|
| SC        | F: ACTGTGGTAGGTGGAGTAC  | 58                         |
|           | R: ACTTGGTGGAGACCTTGT   |                            |
| CYP51     | F: AGATATTGTTGAGAACCTTA| 58                         |
|           | R: CATCAATAGAGCAATCATCT|                            |
| SQLE      | F: ATTTTCGCCCTCATACTTA  | 58                         |
|           | R: AACTGATGCTGAAGGTTGA  |                            |
| ERG6      | F: TGAAACAATCTATGACCTT  | 58                         |
|           | R: GTGATTTGCTGAAGAAGG   |                            |
| ABCB1     | F: CAGCAGTGACGCGCTTAT   | 58                         |
|           | R: ATAGAGTGAAGTTGGAGT   |                            |
| RPA1      | F: ATGGCAACCTTGTCTTCCT  | 58                         |
|           | R: CGTAAAGTGGCGATAGC    |                            |
| ATM       | F: CTCAATGGGTCTGCGGAA   | 58                         |
|           | R: AGGCTTGAGTGATGAAT    |                            |
| RNP       | F: GATTGTGTTGGAGAAACTT  | 58                         |
|           | R: TGCCGATTCTGTGGTAAAC  |                            |
| KRE33     | F: CTATTCCATCCAGTCAA    | 58                         |
|           | R: TGGCCTATGTGCGCTATGA  |                            |
| PheRS     | F: CTTGACGGCAGAACTACTT  | 58                         |
|           | R: GGTCTCGGCTTATGCAGA   |                            |
RESULTS

Identification of *M. canis*

The isolated *M. canis* colony was white, the mycelium on the PDA medium appeared in the form of white wool, and the colony exuded a yellow pigmented secretion (Fig. 1A). After Diff-Quik staining, the fungus was found to be fusiform, thick, rough, thorny, and able to produce large conidia (Fig. 1B). The DNA sequencing results showed that the 18s rDNA sequence of the isolated fungal strain was highly homologous to the 18s rDNA sequence of the *M. canis*. A phylogenetic tree was constructed with similar 18s rDNA sequences (Fig. 2). The results indicated that the isolated fungal strain is closely related to *M. canis* area under the first moment curve. According to the phylogenetic analysis of the morphology and 18s rDNA sequence, the fungal strain was identified as *M. canis*.

Determination of MIC of KTZ

As shown in Fig. 3, KTZ plays a significantly inhibitory role in the growth of *M. canis*. The results showed that the MIC of KTZ against *M. canis* was 1.56 μg/mL.

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Fig. 1. Morphological characteristics of *M. canis*. (A) Colonies of *M. canis* strain on potato dextrose agar plates. (B) Diff-Quik staining of *M. canis* (10 × 40).

Fig. 2. Homology analysis based on partial 18s ribosomal DNA sequences of the *M. canis* and the related microorganisms.

AUMC, area under the first moment curve.

*The DNA sequence of *M. canis* compared here.*
Statistics pertaining to DEGs

As shown in the volcano diagram of DEGs in Fig. 4, there are 779 DEGs, of which 453 DEGs are up-regulated and 326 DEGs are down-regulated. DEGs performed cluster analysis and cluster genes had the same or similar expression patterns. The clustering results of DEGs are shown in Fig. 5. The expression levels of DEGs in the same group were similar. The expression levels of DEGs in different groups differed. The results show that the sample repeatability was good, and that KTZ significantly changed the gene expression of *M. canis.*
GO enrichment analysis

Through GO annotation, 779 DEGs were enriched and analysed from 3 perspectives: biological process, cell composition, and molecular function (Fig. 6).

In the “biological process” category, the subcategories of “biological process” and “cellular process” have the most enriched DEGs. In the category of “cell components”, “cells”, “cell parts”, “organelles”, and “membrane parts” have the most enriched DEGs. In the category of “molecular function”, the 2 most enriched subcategories of DEGs are “binding” and “catalytic activity”.

KEGG annotation and pathway enrichment analysis

As shown in Fig. 7, DEGs are highly enriched in the 3 pathways of RNA polymerase, steroid biosynthesis, and ribosome biogenesis in eukaryotes. Therefore, these 3 pathways were selected for follow-up research.

qRT-PCR identification of DEGs

Ten DEGs, including SC, CYP51, SQLE, ERG6, ABCB1, RPA1, ATM, RNP and KRE33, were screened from 3 pathways of RNA polymerase, steroid biosynthesis, and ribosome biogenesis in eukaryotes (Table 2). The expression levels of DEGs were identified by qRT-PCR. As shown in Fig. 8, DEGs SC, CYP51, SQLE, ERG6, and ABCB1 in the KTZ group were up-regulated ($p < 0.05$), and the expressions of RPA1, ATM, RNP and KRE33 were down-regulated compared
with the control group ($p < 0.05$). The experimental results of qRT-PCR were aligned with the results of transcriptome sequencing.

**DISCUSSION**

The antifungal mechanism of the most commonly used clinical imidazole drugs is very complex. In addition to inhibition of the activity of 14α-demethylase, some imidazole drugs also exert an inhibitory effect on other related enzymes present on the cell membrane [8,14]. For example, when *Candida albicans* is treated with voriconazole, an accumulation of yeast sterols and squalyl alcohol is found [8,15], however, it is unclear whether this result is ascribed to the interaction between voriconazole and some enzymes related to ergosterol synthesis on the cell membrane, or the secondary effect of 14α-demethylase activity after being inhibited. Besides, the mechanism of action of imidazole drugs is also related to the target, such as FLZ and IT, in addition to inhibiting the 14α-demethylase activity of *Cryptococcus neoformans*, it can also inhibit the reduction of obtusifoliol to the corresponding obtusifoliol [16,17]. This effect causes the accumulation of methylated sterol precursors to affect the function of cell membranes [16,17], therefore, the antifungal mechanism of imidazole drugs requires further study. KTZ is commonly used in clinical practice to treat *M. canis* infections. To study the antifungal mechanism of KTZ on *M. canis*, we collected and purified a fungus from dogs with fungal skin diseases. After morphological observation and DNA sequence alignment, the fungus was identified as *M. canis*. 

Fig. 6. GO enrichment analysis of DEGs. The abscissa is the GO classification, the left-hand ordinate is the percentage of the number of genes, that on the right is the number of genes.

GO, gene ontology; DEG, differentially expressed gene.
Based on the experiments of Xiao et al. [18], we used a half MIC of KTZ to treat *M. canis*. This is a stress concentration for *M. canis* that can better reflect the antifungal mechanism of KTZ on *M. canis*. Sequencing results showed that after KTZ treatment, a total of 779 genes were significantly differentially expressed, of which 453 genes were up-regulated and 326 genes were down-regulated. These genes will be the key to explore the antifungal mechanism of KTZ on *M. canis*.

In the results of GO enrichment analysis, the DEGs were most abundant in “biological process”, “cell process”, “cell”, “cell part”, “organelle”, “membrane part”, “binding”, and
“catalytic activity” subcategories. Coincidentally, inhibition of fungal cytochrome P-450-dependent 14α-demethylase changes the permeability of the cell membrane, resulting in the loss of intracellular nutrients and organelles; this is the main antifungal mechanism of KTZ [8-10], therefore, we speculated that the antifungal mechanism of KTZ against *M. canis* was closely related to the presence of these differential genes.

The further to study the role of these differential genes in the antifungal mechanism of KTZ on *M. canis* we screened 3 pathways of RNA polymerase, steroid biosynthesis, and ribosome biogenesis in eukaryotes by KEGG enrichment analysis. The DEGs in these 3 pathways were verified by qRT-PCR. The results of qRT-PCR of the differential genes are consistent with the sequencing results, which show that our sequencing results were correct.

Among the DEGs, CYP51 (sterol 14α-demethylase) is the most conserved cytochrome P450 enzyme in the entire system [19]. In eukaryotes, the P450 enzyme is an enzyme necessary for sterol biosynthesis, and sterol is an essential component of the plasma membrane, which is essential for maintaining the function of fungal cell membranes [20,21]. CYP51 protein is also a target for azole drugs. It demethylates the 14-α position of lanosterol to form ergosterol to change the permeability and fluidity of the cell membrane [21,22]. The up-regulation of CYP51 expression in this experiment indicated that KTZ may play an antifungal role by mediating the P450 enzyme regulated by CYP51.

The other 2 genes related to the cell membrane are squalene monooxygenase (SQLE) and ERG6. The SQLE is a key flavin adenine dinucleotide-dependent enzyme in the biosynthesis pathway of ergosterol and cholesterol, and it is a drug used to inhibit the growth of pathogenic fungi or lower cholesterol levels in potential targets [23]. Zare et al. [24]. found that, after treating *C. albicans* with Te nanoparticles (NPs) (0.2 mg/mL), Te NPs can inhibit SQLE and lead to an increase in the level of SQLE gene expression. This is consistent with our experimental results. Therefore, we speculate that KTZ can inhibit SQLE, so that squalene accumulates to kill *M. canis*. The ERG6 gene encodes the S-adenosylmethionine-dependent sterol C-24 methyltransferase in the ergosterol biosynthetic pathway [25]. Studies have shown that when fungi are resistant to azole drugs, ERG3 and ERG6 may play a role in the biosynthetic pathway of ergosterol [26]. The expression of ERG6 was significantly up-regulated in this experiment, which may be related to the resistance of *M. canis* to KTZ. In
addition, the active efflux of ATP binding cassette (ABC) superfamily proteins is found to be an important mechanism for resistance among azole antifungal drugs [8]. Therefore, we also speculated that the 2 genes of the ABC, ATM and ABCB1, are also involved in the resistance mechanism of *M. canis* to KTZ.

KTZ also affected genes related to the cell wall of *M. canis*. Among differential genes, the expression of Chitinase (SC) gene was significantly up-regulated. SC is an enzyme that can cleave the β-(1,4) glycosidic bond of chitin in the cell wall [27]. Chitin is an important component of the fungal cell wall, and is essential for maintaining the structural integrity of the fungal cell wall and the growth and survival of the fungus [28,29]. Therefore, KTZ may damage the cell wall of *M. canis* by regulating SC.

KTZ not only inhibits those genes related to the cell membrane and cell wall of *M. anis,* but also affects the genes responsible for *M. canis* growth, reproduction and transcription regulation. KRE33 (NAT10) is an N-acetyltransferase and studies have shown that NAT10 may play an important role in cell division by promoting the reorganisation of nucleolus and middle body in the late mitosis and the stabilisation of microtubules [30]. RPA1 is an RNA polymerase, and RNA Pol I is a highly synthetic enzyme that can transcribe rDNA and regulate the growth of eukaryotic cells [31]. RNA polymerases I and III are responsible for most nuclear transcription in actively growing cells, and their activity affects the biosynthetic capacity of the cell [32]. Ribonucleoprotein (RNP) is a protein associated with RNA [33]. It can combine with mRNA and non-coding RNA to form an RNP complex. They can coordinate the production of function-related proteins, so that the activity and stability of many RNAs are regulated after transcription [33,34]. The differential expressions of these genes change significantly, so we speculated that KTZ can inhibit the mitosis of *M. canis* cells by down-regulating the expression of the KRE33 gene and mediating the transcriptional regulation of *M. canis* by regulating RPA1 and RNP.

Our results revealed the potential mechanism of KTZ treatment against *M. canis* through transcriptome sequencing; however, the more detailed aspects of these findings should be clarified in future studies of gene function, such as gene knock-out experiments, so as to better describe the genes selected in this study.

In conclusion, 779 DEGs were screened by transcriptome sequencing. Among them, 453 DEGs were up-regulated and 326 DEGs were down-regulated. The results of GO and KEGG enrichment analysis showed that KTZ could change cell membrane permeability, destroy the cell wall, affect drug resistance, and inhibit mitosis and transcriptional regulation by regulating CYP51, SQL, ERG6, ATM, ABCB1, SC, KER33, RPA1, and RNP in *M. canis*.

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