Selection and evaluation of appropriate reference genes for RT-qPCR based expression analysis in *Candida tropicalis* following azole treatment

Saikat Paul, Shreya Singh, Arunaloke Chakrabarti, Shivaprakash M. Rudramurthy & Anup K Ghosh*

*Candida tropicalis* arises as one of the predominant non-*Candida albicans* Candida (NCAC) species causing invasive candidiasis in Asian countries. A rise in reports of *C. tropicalis* with a parallel increase in fluconazole resistance has also been observed. The genes and underlying pathways associated with azole antifungal resistance in *C. tropicalis* is still not properly understood. The RT-qPCR is the most promising approach for expression analysis of target genes to understand the mechanisms of resistance. The reliability and reproducibility of this technique depend on the selection of suitable reference genes for the normalization in expression study. The present study investigated the expression stability levels of ten genes including ACT1, EF1, GAPDH, PGK1, RDN5.8, RDN18, RDN28, SDHA, TUB1, and UBC13 for their suitability in fluconazole treated/untreated *C. tropicalis*. The stability levels of these genes were examined by the ∆∆CT, ΔCT, Pfaffl methods and five independent software including hkgFinder, geNorm, NormFinder, BestKeeper, and Reffinder software. We report, the EF1 and ACT1 were the most stable reference genes for normalization and can be used for the gene expression analysis in *C. tropicalis*. To the best of our knowledge, our study is the first to select and validate the reference genes in *C. tropicalis* for RT-qPCR based expression analysis.

*Candida tropicalis*, a non-*Candida albicans* Candida (NCAC) resides in human skin, genitourinary, respiratory, and gastrointestinal tracts as a part of the normal microbiota. In Asian countries, *C. tropicalis* has emerged as the predominant NCAC species causing invasive candidiasis (IC), particularly candidemia. Fluconazole is the most common antifungal drug used to treat candidemia due to *C. tropicalis*. The rise in IC due to *C. tropicalis* has been paralleled with an increase in fluconazole resistance, especially in Asian countries. The differential expression of ergosterol biosynthesis pathway genes, ATP-binding cassette (ABC), and major facilitator superfamily (MFS) drug transporters are directly linked with theazole resistance in *C. tropicalis*. Although various studies demonstrated the role of these mechanisms in azole resistant *C. tropicalis*, the principle pathways and regulatory circuits implicated are yet to be elucidated.

Profiling of gene expression is a powerful approach to determine the pattern response to various stimuli including drugs and gives a holistic impression of cellular function in any living cell. Usually, gene expression can be estimated by multiple methods, including RNase protection assay, Northern blotting, real-time quantitative PCR (RT-qPCR), and semi-quantitative reverse-transcription PCR. RT-qPCR has received special attention due to its significantly higher accuracy, sensitivity, and rapidity allowing high throughput results, detection of mRNAs with low-abundance and mRNA copy number measurement. As a result, RT-qPCR platform has been utilized for diverse applications including gene expression analysis.

However, the correctness of the results of RT-qPCR depends on numerous technical and biological factors, including the type of samples, method of sample collection, extraction efficiency of RNA, quality and quantity of RNA input, RNA degradation, cDNA synthesis, PCR efficacy, and errors in pipetting. Additionally, the
sensitivity, reliability, and reproducibility of RT-qPCR based target gene expression measurement depend on the appropriate normalization. Commonly, normalization is performed by using an internal control gene also recognized as housekeeping gene or reference gene. The selection of inappropriate reference genes for RT-qPCR based expression analysis has produced confusing and unreliable results. An appropriate reference gene must be non-regulated, stably expressed, and remain unaltered by experimental and biological conditions. However, no gene with all these characteristics has been identified as yet.

The most frequently utilized reference genes for expression analysis, like 18S and 28S ribosomal RNAs, β-actin, tubulin, and glyceraldehyde 3-phosphate dehydrogenase, have presented variable levels of expression under different conditions in diverse cells and tissues, and are consequently inappropriate for the normalization of RT-qPCR. This suggests the necessity to select and validate the appropriate reference genes which are specific for the type of sample and experimental condition used in different studies.

The present study was performed to examine the expression stability of ten candidate reference genes in 20 resistant and 10 susceptible isolates of C. tropicalis, collected from different clinical specimens including blood, cerebrospinal fluid, pus, and ascitic fluid. The gene expression of the isolates was evaluated in the presence and absence of fluconazole. We examined the stability of these 10 reference genes by utilizing eight different approaches including, ∆∆CT, ΔCT, and Pfaffl methods and by using 5 different software like hkgFinder, geNorm, NormFinder, BestKeeper, and RefFinder. Furthermore, the selected stable reference genes were validated by analyzing the relative expression levels of different pleiotropic azole resistance genes by using the comparative ∆∆CT method.

**Results**

**CT distribution of the reference genes.** Figure 1 representing the CT distributions of 10 candidate reference genes in 60 samples (30 fluconazole treated and 30 untreated control). The instrument generated CT values of the candidate reference genes were ranging from 10.26 to 28.31 (Fig. 1). Four candidate reference genes (EF1, RDN18, RDN28, and GAPDH) presented significantly lower CT values (p < 0.01), indicating a higher abundance of mRNA transcripts. The CT values of RDN18 and RDN28 were uniformly less (~5) in all samples, indicating a higher level of expression. Subsequently, for the stability analysis of RDN18 and RDN28, all the samples (n = 60) were diluted 100 times to increase the CT value up to a detection level (~11). It was difficult to analyse the CT values of RDN18 and RDN28 simultaneously with the other reference genes since undiluted samples were used for their analysis. Therefore, RDN18 and RDN28, could not be utilized as reference genes for the expression analysis of target genes.

**Stability analysis of reference genes in C. tropicalis following fluconazole stimulation by ∆ΔCT, ΔCT and Pfaffl method.** Stability ranking by ∆ΔCT method. To determine the expression stability of the reference genes in fluconazole treated C. tropicalis, the CT values were compared between the untreated control (u) and fluconazole treated (t) cells utilizing the formula: average CT Change = CT(u) - CT(t). Two ribosomal RNA subunits RDN18 and RDN28, EF1, SDHA, UBC13, and GAPDH were the highly stable genes with CT changes < 0.5. Whereas ACT1, PGK1, RDN5.8, and TUB1 were comparatively less stable reference genes with CT changes > 0.5. The stability of RNA expression was validated by comparing with the EF1, as it was found to be both suitable and stable. The ΔCT between EF1 and reference genes was computed by the following
formula: \[ \Delta CT(t) = CT(t \text{ reference}) - CT(t \text{ EF1}) \] and \[ \Delta CT(u) = CT(u \text{ reference}) - CT(u \text{ EF1}) \]. The \[ \Delta \Delta CT(t) \] was calculated by subtracting the \[ \Delta CT \] of untreated cells from the treated cells. Finally, the levels of reference gene expression in the presence of fluconazole was calculated by transforming the \[ \Delta \Delta CT \] into \[ 2^{-\Delta \Delta CT} \] value. The computed \[ \Delta \Delta CT(t) \] and \[ 2^{-\Delta \Delta CT} \] results of the 10 candidate reference genes in fluconazole treated samples are provided in Table 1. The \[ 2^{-\Delta \Delta CT} \] values indicate that \( \text{EF1} \), \( \text{SDHA} \), \( \text{RDN18} \), \( \text{RDN28} \), \( \text{UBC13} \), and \( \text{GAPDH} \) were the most stable, while \( \text{ACT1} \), \( \text{PGK1} \), \( \text{RDN5.8} \), and \( \text{TUB1} \) were comparatively less stable.

**Analysis by \( \Delta CT \) method.** The stability ranking of the reference genes was also analyzed by the \( \Delta CT \) method. Among the 10 reference genes examined in this study, \( \text{RDN18}, \text{ACT1}, \text{and RDN28} \) were the most stable reference genes, while \( \text{UBC13}, \text{PGK1}, \text{and TUB1} \) were the least stable genes (Fig. 2). However, a difference in the stability ranking of the most stable reference genes selected by \( \Delta \Delta CT \) and \( \Delta CT \) methods was seen.

**Analysis by Pfaffl method.** The expression stability of the reference genes was measured by using the Pfaffl method. According to the ranking order of the reference genes, \( \text{EF1}, \text{SDHA} \) and \( \text{RDN18} \) were the most stable, whereas \( \text{PGK1}, \text{RDN5.8} \), and \( \text{TUB1} \) were the least stable reference genes (Table 2). The stability ranking of reference genes by the Pfaffl and \( \Delta \Delta CT \) methods were exactly the same and were different from the \( \Delta CT \) method (Table 3).

**Stability ranking of reference genes using five different software.** Five independent software including hkgFinder, geNorm, Norm-Finder, BestKeeper, and web-based RefFinder software were utilized to calculate the stability levels of the genes tested. Each software utilizes a considerably different algorithm to assess...
the stability of the reference genes. Results obtained from these five distinct approaches were used to select the most stable reference genes.

**hkgFinder analysis.** The hkgFinder software selects the most stable reference genes by grading the candidate reference genes with respect to their standard deviation (SD) and fold changes (FC) (Table 4). Out of the 10 candidate genes, the SDs were between 0.58 to 1.64, and the FCs between 1.1 to 1.7. According to hkgFinder, the most stable candidate reference genes identified were EF1, RDN28, and RDN28.

**geNorm analysis.** The geNorm software measures the stability levels of the candidate genes by computing the ‘M’ stability score. A lower an M score indicates higher stability and the default limit of 1.5 is recommended as cut off. All the reference genes had an M score < 1.5 and the geNorm software selected RDN18, RDN28, EF1, and ACT1 as the most stable reference genes (Fig. 3).

The geNorm software also suggests whether a combination of reference genes is needed or not. Each normalization factor (NF) computed the geometric mean values of two reference genes and calculated their pairwise variability (V value). A combination of reference genes is not recommended at V value < 0.15. In this study, all combinations of genes showed V value < 0.15, indicating no need for combining reference genes for normalization (Fig. 4 and Supplementary Table S4).

**NormFinder analysis.** The NormFinder software analysis grades the reference gene depending upon the stability score, calculated from the intergroup and intragroup expression variability. Although the best reference genes selected by the hkgFinder and geNorm programs were similar (RDN18, RDN28, and EF1) they were considerably different from NormFinder (ACT1, RDN18, and GAPDH) (Fig. 5 and Table 3).

**BestKeeper analysis.** BestKeeper program allows for a comparative measurement between different reference genes. The analysis of 10 reference genes exhibited a significantly higher correlation of 0.803 ≤ r ≤ 0.932 among their levels of expressions and the BestKeeper index (r), however, the best correlations were observed for ACT1, RDN18, and PGK1 (Fig. 6).

The BestKeeper software also determined the expression stability by computing both the standard deviation (SD) and as well as the coefficient of variance (CV) of the mean CT values. Out of 10 candidate genes, eight showed SD within the recommended range [0.5 < SD(±CT) ≤ 1.00]. EF1 showed the lowest SD (0.37) and CV

---

**Table 4.** BestKeeper software base descriptive statistical analysis of reference genes.

| Gene   | ACT1 | GAPDH | PGK1 | RDN 5.8 | RDN 18 | RDN 28 | SDHA | TUB1 | UBC 13 | EF1 |
|--------|------|-------|------|---------|--------|--------|------|------|--------|-----|
| N      | 60   | 60    | 60   | 60      | 60     | 60     | 60   | 60   | 60     | 60  |
| GM (CT)| 15.14| 11.45 | 18.11| 12.23   | 11.19  | 11.60  | 11.74| 23.69| 21.02  | 11.34|
| AM (CT)| 15.17| 11.48 | 18.17| 12.28   | 11.22  | 11.62  | 17.26| 21.05| 11.35  |
| Min (CT)|13.74| 10.26 | 15.95| 10.60   | 10.45  | 10.90  | 16.12| 19.20| 10.60  |
| Max (CT)|19.32| 14.47 | 23.68| 17.55   | 14.56  | 15.31  | 21.12| 24.32| 14.23  |
| SD (±CT)|0.62 | 0.62  | 1.23 | 0.83    | 0.49   | 0.47   | 0.63 | 1.37 | 0.90   | 0.37|
| CV (% CT)|4.11 | 5.41  | 6.76 | 6.73    | 4.39   | 4.04   | 3.64 | 5.76 | 4.26   | 3.28|
| Min (x-fold)|−2.64| 2.28  | 4.46 | 3.09    | −1.67  | −1.63  | −2.17| −7.01| −3.54  | −1.67|
| Max (x-fold)|18.13| 8.13  | 47.64| 39.96   | 10.32  | 13.05  | 14.73| 24.59| 9.83   | 7.42|
| SD (±x-fold)|1.54 | 1.54  | 2.34 | 1.77    | 1.41   | 1.39   | 1.55 | 2.58 | 1.86   | 1.29|

---

**Figure 3.** Expression stability values (M) of ten reference genes measured by the geNorm program. Candidate reference genes are ranked from left to right according to their increasing of stability (declining M values).
indicating higher stability, while PGK1 and TUB1 were not found stable as their SD (1.23 and 1.37 respectively) and CV (6.76 and 5.76 respectively) were higher (Table 5). The reference genes (RDN18, RDN28, and EF1) selected by BestKeeper, geNorm, and hkgFinder were similar while slight discordance with NormFinder was seen (Table 3).

**RefFinder analysis.** The RefFinder was utilized for the final ranking of the reference genes tested. RefFinder, the web-based tool analyzed the data by integrating NormFinder, BestKeeper, and geNorm for final grading of the reference genes. RDN18, RDN28, ACT1, and EF1 were selected as the best reference genes under every experimental setup (Fig. 7).

Overall RDN18, RDN28, EF1, and ACT1 were the most stable genes. As the amplification efficiency of RDN18 and RDN28 was very high, they were excluded. Further, EF1, ACT1 and the next most stable genes GAPDH, and SDHA were evaluated. (Table 3).
Evaluation of selected reference genes. Using EF1, ACT1, GAPDH, and SDHA as internal control, the inducible expression of azole resistance related genes, CDR1, CDR2, MDR1, ERG1, ERG3, and ERG11 was examined among the resistant isolates. The presence of fluconazole, noticeably increased the expression levels of all the genes tested when normalized with EF1 (2.1 to 9.7 fold) and ACT1 (2.1 to 7.1 fold). The expression levels of azole resistance related genes were comparatively lower when normalized with GAPDH (1.2 to 5.8 fold) and SDHA genes (1.1 to 3.3 fold). However, this variation was not significant (p > 0.05) indicating that any of these genes may be utilized for normalization in inducible expression analysis of resistance related genes (Fig. 8).

Table 5. Ranking of C. tropicalis reference gene with respect to expression stability as analysed by six different approaches.

| Ranking | \(2^{-\Delta\Delta CT}\) | \(\Delta CT\) | Pfaffl | hkgFinder | geNorm | NormFinder | BestKeeper | RefFinder |
|---------|----------------|--------|-------|----------|--------|------------|------------|----------|
| 1       | EF1            | RDN18  | EF1   | EF1      | RDN18  | ACT1       | ACT1       | RDN18    |
| 2       | SDHA           | ACT1   | SDHA  | RDN28    | RDN18  | RDN18      | RDN18      | RDN28    |
| 3       | RDN18          | RDN28  | RDN18 | EF1      | GAPDH  | PGK1       | ACT1       | RDN28    |
| 4       | RDN28          | GAPDH  | RDN28 | SDHA     | ACT1   | RDN28      | RDN28      | EF1      |
| 5       | UBC13          | EF1    | UBC13 | GAPDH    | GAPDH  | SDHA       | RDN5.8     | GAPDH    |
| 6       | GAPDH          | SDHA   | GAPDH | ACT1     | SDHA   | EF1        | EF1        | SDHA     |
| 7       | ACT1           | RDN5.8 | ACT1  | UBC13    | RDN5.8 | UBC13      | UBC13      | RDN5.9   |
| 8       | PGK1           | UBC13  | PGK1  | RDN5.8   | PKG1   | PKG1       | PKG1       | UBC13    |
| 9       | RDN5.8         | PKG1   | RDN5.8| PKG1     | PGK1   | PKG1       | UBC13      | PKG1     |
| 10      | TUB1           | TUB1   | TUB1  | TUB1     | TUB1   | TUB1       | TUB1       | TUB1     |

Figure 7. Ranking of reference genes by using the web-based RefFinder tool.

Discussion
Appropriate normalization strategies are crucial for the correction of variability in the multistep process of gene expression analysis. Most RT-qPCR experiments are performed by using a single reference gene for normalization. A study by Vandesompele et al. highlighted that ACT1, GAPDH, RDN18, and RDN28 are the most frequently utilized single reference genes for normalization. However, only a few studies have paid attention to the appropriate validation of candidate reference genes used in RT-qPCR based gene expression analysis.

In our study, the expression stability analysis of ten reference genes was performed to select the best internal controls for normalization. The expression levels of all the candidate genes were found to be differentially stable and suitable in fluconazole treated C. tropicalis isolates. To our knowledge, this is the first study to select and validate the appropriate reference genes for expression analysis in clinical isolates of C. tropicalis.

We used eight different methods including \(\Delta\Delta CT\), \(\Delta CT\), Pfaffl, hkgFinder, geNorm, NormFinder, BestKeeper, and web-based RefFinder for stability analysis. The most stable reference gene identified was variable among these eight methods, possibly due to the differences in their analytical algorithms. Similarly, stable expression of RDN18 and RDN28 was seen in the present study while the clear unsuitability of RDN5.8 as an internal control was noted. The use of RDN18 and RDN28 has been recommended by several studies as an internal control for mRNA quantification. However, since the transcript levels of RDN18 and RDN28 were very high (CT~5) and substantial sample dilution was required, it was difficult to correctly deduce the baseline values in expression analysis. As a result, RDN18 and RDN28 genes were excluded as reference genes despite their high stability.

Seven other reference genes with diverse functions were selected for further investigation in the present study. These genes could be categorized into the following classes: transcription-related genes (EF1), glycolytic enzymes (GAPDH, and PGK1), citric acid cycle enzyme (SDHA), cytoskeleton-related genes (ACT1, and TUB1), and Ubiquitin-conjugating enzyme (UBC13). Multiple studies had been performed using these genes as internal controls. The present study demonstrated that EF1 was one of the most stable genes, which is contradictory to
to the findings of Anita et al.\textsuperscript{10}. \textit{ACT1}, \textit{TUB1}, and \textit{SDHA} are also commonly utilized reference genes in stability analysis and target gene expression studies.\textsuperscript{10,17,34–36,38} A study by Li et al. clearly demonstrated the unsuitability of \textit{ACT1}, \textit{TUB1}, and \textit{SDHA} as reference genes for inducible expression analysis in \textit{C. glabrata} cells treated with fluconazole\textsuperscript{17}. In contrast, these genes were recommended as the most suitable and stable reference genes in \textit{Microsporum canis} under various experimental conditions\textsuperscript{10}. Such heterogeneous results may be due to the inherent biological characteristics of the different fungal isolates and/or variation in the test conditions. In the present study, the stability and suitability of \textit{ACT1}, \textit{SDHA}, and \textit{EF1} were comparable making them all appropriate as reference genes for expression analysis in \textit{C. tropicalis}. However, \textit{TUB1} was found to be comparatively less stable. Additionally, \textit{UBC13} was found to unsuitable in the present study, which is in contrast to the findings of Li et al.\textsuperscript{17}. The \textit{PGK1} and \textit{GAPDH} genes play a significant role in the glycolytic pathway and variable degrees of expression stability have been reported by several studies\textsuperscript{10,17,21,23,28,33–36,38,39}. This variation may be due to the difference in organisms selected for analysis and different experimental conditions utilized. \textit{PGK1} and \textit{GAPDH} had been shown to be co-regulated in a previous study\textsuperscript{40}. However, such co-regulation was not evident in the present study and \textit{GAPDH} was more stable compared to \textit{PGK1}. The validation of \textit{EF1}, \textit{ACT1}, \textit{GAPDH}, and \textit{SDHA} as an internal control in azole resistance gene expression analysis was performed which confirmed either of these genes could be used as a potential reference gene. Thus, the arbitrary selection of a reference gene must be avoided and validation of internal controls across different experimental setup is essential.

Even though normalizing with a single reference gene is simple to use and widely accepted, some researchers have recommended the utilization of more than one reference gene for normalization\textsuperscript{17,21,23,41}. This is the pragmatic approach to ensure correct normalization which is particularly essential when dealing with fine measurements. Vandesompele \textit{et al.} documented that normalization using a single reference gene may lead to erroneous results and additional reference genes may be required\textsuperscript{21}. Additionally, the stability analysis of multiple reference genes is not always possible, due to the less sample availability and significantly higher running cost. Furthermore, by using different combinations of multiple reference genes, the inter-experiment variability in result interpretation may increase. Using geNorm analysis, we found that a single reference gene was sufficient for providing accurate normalization and the combination of the reference gene is not required.

**Conclusion**

In conclusion, the present study is the first to select and validate the reference genes in \textit{C. tropicalis} for RT-qPCR based expression analysis. Our study highlights that, evaluation of the most appropriate internal control is an important prerequisite for RT-qPCR based expression analysis in different experimental models. The present study may also give preliminary knowledge for the assessment of candidate genes for expression studies in different \textit{Candida} species and in diverse experimental conditions.
**Materials and Methods**

**Isolates and growth conditions.** The present study was conducted by following the Minimum Information for Publication of Quantitative Real-Time PCR Experiments (MIQE) guidelines. Twenty fluconazole-resistant (16–256 μg/mL) and 10 susceptible (0.5–1 μg/mL) isolates of *C. tropicalis* from invasive candidiasis were used in this study. Of the 30 isolates collected from the patients, 26 from the blood, 2 from cerebrospinal fluid, and one each from pus and ascitic fluid (Supplementary Table S1). Informed consent was taken from each enrolled patient or a parent/guardian if the patient is under 18 following our institute protocol. All the experiments used in this study were performed as per the guidelines and regulations approved by the institutional ethics committee of the Postgraduate Institute of Medical Education and Research (PGIMER), Chandigarh, India. All these isolates were also used in our previous studies and their details MICs are presented in Supplementary Table S1. A total of 60 samples (30 fluconazole treated and 30 untreated controls) were used for stability analysis. The confirmation of identification was done by both PCR sequencing of the ITS region and by using MALDI-TOF MS (Bruker).

**Extraction of RNA and cDNA synthesis.** Total RNA from *C. tropicalis* isolates was extracted at the logarithmic phase using TRizol reagent (Invitrogen, USA) as per the manufacturer’s protocol. Both the quantity and quality of the extracted total RNA was analysed by determining the absorbance (A260/A280) using a spectrophotometer (NanoDrop 2000/2000c, Thermo Scientific, USA). The RNA samples with 260/280 ratio of 1.85 to 2.06 were used in the present study. The integrity of the RNA samples was further examined by running 1% denaturing agarose gel. An RNAase-free DNAse treatment (Qiagen, Germany) was given to each RNA preparation. One microgram of total RNA input was used in a 20 μL reaction volume. The PCR amplification was executed by using the standard protocol with the Eppendorf 5331 MasterCycler. The PCR amplification was executed by using the standard protocol with the Eppendorf 5331 MasterCycler.

**Candidate gene selection and primer design.** Ten candidate genes including *ACT1, EF1α, GAPDH, PGK1, RDN5.8, RDN18, RDN28, SDHA, TUB1, and UBC13* were examined for expression stability (Table 6). For the validation of reference genes 6 target genes [ABC transporter genes (CDR1 and CDR2), Multi drug resistance gene (MDR1), Squalene epoxidase (ERG1), Δ5,6-desaturase (ERG3), Lanosterol C14α demethylase (ERG11)] related to azole resistance were also studied (Supplementary Table S2). The coding sequences of these candidate genes were obtained from NCBI (www.ncbi.nlm.nih.gov) and Candida Genome Database (www.candidagenome.org) (Table 6 and S2). Sequences of the selected genes were used to synthesize the primers by using the web-based Primer-Blast tool in NCBI and the quality of each primer was checked by using the online Sequence Manipulation Suite tool (www.bioinformatics.org) (Table 6 and S3). The binding efficiency of the synthesized primers was determined and the standardized optimum primer annealing temperature was 59°C.

**RT-qPCR analysis.** RT-qPCR amplification of reference genes and target genes was performed by using Light Cycler 480 (Roche, Switzerland). The 10 μL reaction mixture contained 5 μL of PowerUp SYBR Green Master Mix.
(Thermo Fisher Scientific, United States), 0.25 μL each primer (10 pmol), 1 μL cDNA and 3.5 μL nuclease-free ultrapure water. Amplification of cDNA templates was executed by employing the following conditions: starting denaturation cDNA templates at 95 °C for 1 min, then repetitions 45 cycles of denaturation at 94 °C for 10 seconds, annealing at 59 °C for 10 seconds, and extension at 72 °C for 10 seconds. Finally, the melting curve assessment was completed by applying the setup of denaturation at 95 °C for 5 seconds, annealing at 59 °C for 1 min and 97 °C for 15 seconds. The ‘CT’ or threshold cycle is the number of the cycle at which the reporter dye used in the PCR reaction crossed the software-designated threshold, which was automatically calculated by the Light Cycler 480 System Software (Roche, Switzerland) versions 1.5.

The calculated amplification efficiencies (E) for all the candidate genes used in this study were between 93.3–99.7% (Supplementary Table S5). The standard curves for the 16 genes were constructed by three fold serial dilutions and linear correlation coefficients (R²: 0.994–1.00) were observed (Supplementary Fig. S1). Each primer sets used in this study generated a single peak indicating the production of a single product (Supplementary Fig. S2). For each RT-qPCR experiment, two negative controls including one containing all the components without cDNA and another without primers were assessed simultaneously.

**Stability analysis of constitutively expressed genes.** CT values from the RT-qPCR instrument were used for the stability analysis of reference genes to select the best genes for inducible expression analysis of pleiotropic target genes. The basal or constitutive expression level of reference genes was determined by comparing the fold changes with respect to a stable reference gene as a comparator by using ΔΔCT27, ΔCT39, Pfaffl37 approach. Five different software: hkgFinder17, geNorm14, NormFinder30, BestKeeper31, and RefFinder32 were used for the stability assessment of reference genes. The hkgFinder algorithm calculates the standard deviation (SD) of CT values obtained from both azole-untreated and treated cells of C. tropicalis and also calculates the fold changes among both phenotypes. The smallest SD of reference genes indicated the most stable reference gene. Another software, geNorm calculates both the stability value (M) and a pairwise variation (V). Both these analysis are used to examine the stability of any reference gene and to assess whether the combination of reference genes is required or not. The NormFinder software calculates the stability values depending on the intergroup and intragroup variability in the expression of different reference genes. The BestKeeper software determined the pairwise correlation to rule out the stability of a reference gene with the BestKeeper Index, that is basically indicating the geometric mean (GM) of the acquired CT values. The BestKeeper algorithm commonly computes both the coefficients of variance CV(%)CT and standard deviation SD(±CT) for all the candidate reference genes. RefFinder is a web-based platform was utilized to assess and screen the candidate genes for stability ranking. It incorporates the most commonly used programs including BestKeeper, NormFinder, and geNorm to analyse and rank the candidate reference genes. The recommended guidelines for each software package was followed by entering the raw RT-qPCR data obtained as an output from the instrument, and the results were analysed accordingly17,47. For the validation of stable reference genes, the inducible overexpression expression of the resistance related genes was measured among the resistant isolates with respect to untreated control by utilizing the ΔΔCT method27.

Published: 29 October 2019; Accepted: 12 December 2019;
Published online: 06 February 2020

**References**

1. Basu, S., Gugnani, H. C., Joshi, S. & Gupta, N. Distribution of Candida species in different clinical sources in Delhi, India, and proteinase and phospholipase activity of Candida albicans isolates. Rev. Iberoam. Microl. 20, 137–40 (2003).
2. Oksuz, S. et al. Distribution and phospholipase activity in different Candida species isolated from anatomically distinct sites of healthy adults. Jpn. J. Infect. Dis. 60, 280–283 (2007).
3. Negri, M. & Martins, M. Examination of Potential Virulence Factors of Candida tropicalis Clinical Isolates From Hospitalized Patients. Mycopathologia 169, 175–182 (2010).
4. Chakrabarti, A. et al. Incidence, characteristics and outcome of ICU-acquired candidemia in India. Intensive Care Med. 41, 285–295 (2014).
5. Wu, P. F. et al. Epidemiology and antifungal susceptibility of candidemia isolates of non-albicans Candida species from cancer patients. Emerg. Microbes Infect. 6, e87 (2017).
6. Fan, X. et al. Notable Increasing Trend in Azole Non-susceptible Candida tropicalis Causing Invasive Candidiasis in China (August 2009 to July 2014): Molecular Epidemiology and Clinical Azole Consumption. Fron. Microbiol. 8, 464 (2017).
7. Fan, X. et al. Molecular mechanisms of azole resistance in Candida tropicalis isolates causing invasive candidiasis in China. Clin. Microbiol. Infect. 25, 885–891 (2019).
8. Jiang, C. et al. Mechanisms of azole resistance in 52 clinical isolates of Candida tropicalis in China. J. Antimicrob. Chemother. 68, 778–785 (2013).
9. Choi, M. J. et al. Resistance Mechanisms and Clinical Features of Flucytosine-Nonsusceptible Candida tropicalis Isolates Compared with Fluconazole-Less-Susceptible Isolates. Antimicrob. Agents Chemother. 60, 3653–3661 (2016).
10. Giesielska, A. & Stączek, P. Selection and validation of reference genes for qRT-PCR analysis of gene expression in Microsporum canis growing under different adhesion-inducing conditions. Sci. Rep. 8, 1197 (2018).
11. VanGuilder, H. D., Vrana, K. E. & Freeman, W. M. Twenty-five years of quantitative PCR for gene expression analysis. Biotechniques 44, 619–626 (2008).
12. Bustin, S. Absolute quantification of mRNA using real-time reverse transcription polymerase chain reaction assays. J. Mol. Endocrinol. 25, 169–193 (2000).
13. Giulietti, A. et al. An overview of real-time quantitative PCR: Applications to quantify cytokine gene expression. Methods 25, 386–401 (2001).
14. Bustin, S. A., Benes, V., Nolan, T. & Pfaffl, M. W. Quantitative real-time RT-PCR - A perspective. J. Mol. Endocrinol. 34, 597–601 (2005).
15. Kubista, M. et al. The real-time polymerase chain reaction. Mol. Aspects Med. 27, 95–125 (2006).
16. Wong, M. L. & Medrano, J. F. Real-time PCR for mRNA quantitation. Biotechniques 39, 75–85 (2005).
17. Li, Q., Skinner, J. & Bennett, J. E. Evaluation of reference genes for real-time quantitative PCR studies in Candida glabrata following azole treatment. BMC Mol. Biol. 13, 22 (2012).
22. Schmittgen, T. D. & Zakrajsek, B. A. Effect of experimental treatment on housekeeping gene expression: validation by real-time, quantitative RT-PCR. *J. Biochem. Biophys. Methods* **46**, 69–81 (2000).

23. Thellin, O. et al. Housekeeping genes as internal standards: use and limits. *J. Biotechnol.* **75**, 291–295 (1999).

24. Selvey, S. et al. β-Actin—an unsuitable internal control for RT-PCR. *Mol. Cell. Probes* **15**, 307–311 (2001).

25. Clarke, E. M., Divjak, M., Bailey, M. J. & Walters, E. H. β-actin and GAPDH housekeeping gene expression in asomatic airways is variable and not suitable for normalising mRNA levels. *Thorax* **57**, 765–770 (2002).

26. Zhong, H. & Simons, J. W. Direct Comparison of GAPDH, β-Actin, Cyclophilin, and 28S rRNA as Internal Standards for Quantifying RNA Levels under Hypoxia. *Biochem. Biophys. Res. Commun.* **259**, 523–526 (1999).

27. Schmittgen, T. D. & Livak, K. J. Analyzing real-time PCR data by the comparative C(T) method. *Nat. Protoc.* **3**, 1101–8 (2008).

28. Silver, N., Best, S., Jiang, J. & Thein, S. L. Selection of housekeeping genes for gene expression studies in human reticulocytes using real-time PCR. *BMC Mol. Biol.* **7**, 33 (2006).

29. Pfaffl, M. W. A new mathematical model for relative quantification in real-time RT-PCR. *Nucleic Acids Res.* **29**, e45 (2001).

30. Andersen, C. L., Jensen, J. L. & Orntoft, T. F. Normalization of Real-Time Quantitative Reverse Transcription-PCR Data: A Model-Based Variance Estimation Approach to Identify Genes Suited for Normalization, Applied to Bladder and Colon Cancer Data Sets. *Cancer Res.* **64**, 5245–5250 (2004).

31. Pfaffl, M. W., Tichopad, A., Prgomet, C. & Neuviants, T. P. Determination of stable housekeeping genes, differentially regulated target genes and sample integrity: BestKeeper – Excel-based tool using pair-wise correlations. *Biotechnol. Lett.* **26**, 509–515 (2004).

32. Xie, F., Xiao, P., Chen, D., Xu, L. & Zhang, B. miRDeepFinder: a miRNA analysis tool for deep sequencing of plant small RNAs. *Plant Mol. Biol.* **80**, 75–84 (2012).

33. Nakayama, T. et al. Assessment of suitable reference genes for RT-qPCR studies in chronic rhinosinusitis. *Sci. Rep.* **25**, 1568 (2018).

34. Song, L., Li, T., Fan, L., Shen, X.-Y. & Hou, C.-L. Identification and Evaluation of Reliable Reference Genes in the Medicinal Fungus Shiraia bambusambica. *Curr. Microbiol.* **72**, 444–449 (2016).

35. St-Pierre, J., Grégoire, J. C. & Vaillancourt, C. A simple method to assess group difference in RT-qPCR reference gene selection using GeNorm: The case of the placental sex. *Sci. Rep.* **7**, 16923 (2017).

36. Qian, J. et al. Selection and Evaluation of Appropriate Reference Genes for RT-qPCR Normalization of Volvariella volvacea Gene Expression under Different Conditions. *Biomed Res. Int.* **2018**, 1–12 (2018).

37. Huis, R., Hawkins, S. & Neutelings, G. Selection of reference genes for quantitative gene expression normalization in flax (Linum usitatissimum L.). *BMC Plant Biol.* **10**, 71 (2010).

38. Zampieri, D., Nora, L. C., Basso, V., Camassola, M. & Dillon, A. J. P. Validation of reference genes in *Penicillium echinulatum* to enable gene expression study using real-time quantitative RT-PCR. *Curr. Genet.* **60**, 231–236 (2014).

39. Sinha, P. et al. Matrix-assisted laser desorption ionization time-of-flight mass spectrometry for the rapid identification of yeasts based rapid detection of fluconazole resistance in *Candida tropicalis*. *Clin. Microbiol. Infect.* **21**, 32–42 (2015).

40. Anderson, L. E., Gatla, N. & Carol, A. A. Enzyme co-localization in pea leaf chloroplasts: glyceraldehyde-3-P dehydrogenase, triose-P isomerase, aldolase and sedoheptulose bisphosphatase. *Photosynth. Res.* **83**, 317–328 (2005).

41. Teste, M.-A., Duquenne, M., François, J. M. & Parrou, J.-L. Validation of reference genes for quantitative expression analysis by real-time RT-PCR in *Saccharomyces cerevisiae*. *BMC Mol. Biol.* **10**, 99 (2009).

42. Bustin, S. A. et al. The MIQE Guidelines: Minimum Information for Publication of Quantitative Real-Time PCR Experiments. *Clin. Chem.* **55**, 611–622 (2009).

43. Paul, S. et al. Rapid detection of fluconazole resistance in *Candida tropicalis* by MALDI-TOF MS. *Med. Mycol.* **56**, 231–241 (2018).

44. Paul, S., Singh, S., Chakrabarti, A., Rudramurthy, S. M. & Ghosh, A. K. Stable isotope labelling: an approach for MALDI-TOF MS-based rapid detection of fluconazole resistance in *Candida tropicalis*. *J. Antimicrob. Chemother.* **74**, 1269–1276 (2019).

45. Ghosh, A. K. et al. Matrix-assisted laser desorption ionization time-of-flight mass spectrometry for the rapid identification of yeasts causing bloodstream infections. *Clin. Microbiol. Infect.* **21**, 372–8 (2015).

46. White, T. J., Bruns, T., Lee, S. & Taylor, J. Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. *PCR Protocols: A Guide to Methods and Applications* 315–322 (1990).

47. Nails, H., Coenye, T., Van Nieuwerburgh, F., Deforce, D. & Nelis, H. J. Development and evaluation of different normalization strategies for gene expression studies in *Candida albicans* biofilms by real-time PCR. *BMC Mol. Biol.* **7**, 1–9 (2006).

Acknowledgements
The authors duly acknowledge the Indian Council of Medical Research (ICMR), Government of India for financial supports.

Author contributions
S.P. designed the study, conducted the experiments, acquired, analysed and interpreted the data and wrote the manuscript. S.S., A.C. and S.M.R. revised it critically for important intellectual content. A.K.G. designed the study, analysed, and interpreted the results and revised it critically. All the authors have approved the final version of the article.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-58744-7.

Correspondence and requests for materials should be addressed to A.K.G.

Reprints and permissions information is available at www.nature.com/reprints.
