Dynamic Transcript Profiling of *Candida albicans* Infection in Zebrafish: A Pathogen-Host Interaction Study

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**Abstract**

*Candida albicans* is responsible for a number of life-threatening infections and causes considerable morbidity and mortality in immunocompromised patients. Previous studies of *C. albicans* pathogenesis have suggested several steps must occur before virulent infection, including early adhesion, invasion, and late tissue damage. However, the mechanism that triggers *C. albicans* transformation from yeast to hyphae form during infection has yet to be fully elucidated. This study used a systems biology approach to investigate *C. albicans* infection in zebrafish. The surviving fish were sampled at different post-infection time points to obtain time-lapsed, genome-wide transcriptomic data from both organisms, which were accompanied with in sync histological analyses. Principal component analysis (PCA) was used to analyze the dynamic gene expression profiles of significant variations in both *C. albicans* and zebrafish. The results categorized *C. albicans* infection into three progressing phases: adhesion, invasion, and damage. Such findings were highly supported by the corresponding histological analysis. Furthermore, the dynamic interspecies transcript profiling revealed that *C. albicans* activated its filamentous formation during invasion and the iron scavenging functions during the damage phases, whereas zebrafish ceased its iron homeostasis function following massive hemorrhage during the later stages of infection. Most of the immune related genes were expressed as the infection progressed from invasion to the damage phase. Such global, interspecies evidence of virulence-immune and iron competition dynamics during *C. albicans* infection could be crucial in understanding control fungal pathogenesis.

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**Introduction**

*Candida albicans* is a major fungal human pathogen [1,2]. Although *C. albicans* exists as a commensal organism on the cutaneous and mucosal surfaces of healthy individuals, it is also an opportunistic pathogen that can cause systemic and chronic infections. In immunocompromised patients, *C. albicans* is responsible for a number of life-threatening infections, causing considerable morbidity and mortality. Prior research has identified several *C. albicans* virulence factors, including yeast-to-hyphae morphogenesis, the expression of cell surface adhesins, and extracellular hydrolytic activities [3–5]. However, the mechanism that triggers *C. albicans* to transform from yeast to hyphae has yet to be fully elucidated [6].

Previous studies of *C. albicans* pathogenesis have suggested several steps that may lead to virulent infection, including early adhesion, invasion, and late tissue damage [7–9]. *C. albicans* may first colonize the host’s epithelial surface and persist on expanding, leading to invasion and tissue damage. In the invasion stage, morphogenesis plays a significant role that involves cell transit from ovoid yeast (blastospore) to a filamentous (pseudohyphae and hyphae) form. Finally, the fungus penetrates deep into tissues and organs, causing severe damage and possible host fatality. The underlying molecular events for such a progression are expected to be regulated by the pathogen-host interaction.

The interaction between *C. albicans* and the host during infection is a complex and dynamic process. *C. albicans* utilizes different strategies to cope with the host’s environmental cues for morphological change, proliferation, and escape from the host’s immune defenses. The environmental conditions within the host include nutrient and chemical factors, such as the availability of metal ions and amino acids, reactive oxygen and nitrogen species, host-secreted antimicrobial peptides, antifungal drugs used in therapy, or changes in temperature and pH [10]. Several genomewide studies have investigated these complex pathogen-host interactions. They have typically utilized a cell line-based system...
to represent the host environment within an organ or tissue. These approaches include *C. albicans* infections in oral epithelial cells [8], macrophages [11], neutrophils [12], dendritic cells [13], THP-1 human mononuclear cells [14], vascular endothelial cells [15], and human blood [16]. Moreover, there are some studies on *C. albicans* infections in living hosts, e.g. mice [17,18] and rabbits [19]. However, none of these prior studies provided the whole genome response of a living host during *C. albicans* infection.

Recently, researchers have demonstrated that zebrafish (*Danio rerio*) can be a useful model for invasive fungal research. Chao et al. 2010 [20] performed two-step homogenization/mRNA extraction of a whole zebrafish infected with *C. albicans* could offer a pool of gene transcripts from both the host and the pathogen, enabling the individual estimation of specific gene expression profiles using sequence-targeted probes derived from individual genomes. Brothers et al. 2011 [21] also used zebrafish to study candidemia and that revealed how the lethal disease takes effect in zebrafish and found NADPH oxidase is required for regulation *C. albicans* filamentous formation. However, those aforementioned studies did not provide genome-wide transcript profiling data to uncover the comprehensive dynamic interaction in either the host or the pathogen. Microarray dataset of fungal infection surviving zebrafish at different post-infection time points was provided by our research group (NCBI Gene Expression Omnibus Database (GSE32119)). Kuo et al. 2013 [22] used this microarray data set to construct infection-related protein-protein interaction (PPI) network and intracellular PPI networks of the two species, the division of stages was based on histological analysis. In this present work, the same dataset was analyzed by a different method. This principal component analysis-based approach allows us to identify significant expression profiles, and classify infection stages based on transcript profiling of both zebrafish host and fungal pathogen that are consistent with the histological observations. The results also indicated that progression of infection and emergence of virulence is highly correlated with iron competition between the host and the fungal pathogen.

**Results and Discussion**

**Dynamic gene expression analysis of *C. albicans* infection in zebrafish**

Time-course gene expression analyses in both *C. albicans* and zebrafish elucidated the molecular basis of *C. albicans* infection within a living host. A total of 6,202 genes were found to exist in the expression profile of *C. albicans*. One-way ANOVA detected significant variations across nine time points. Figure S1A illustrates the numbers of significant genes identified at different *p*-values. Using a *p*-value of ≤0.01 as the cut-off threshold, the data filter procedures selected a total of 4,827 significant genes for *C. albicans*. As mentioned previously, hyphal morphogenesis is an important virulence factor in *C. albicans*. As a validation to the proposed approach, the present study’s findings identified all key genes related to hyphal formation [10], except *NRG1*. Applying a more stringent criterion of a *p*-value less than 0.01/6202 = 1.61×10^-6 (with Bonferroni correction), the target gene pool was further reduced to total of 1,307 significant genes (Table S1). Gene ontology (GO) analysis of these 1,307 significant genes, based on the *Candida* Genome Database (http://www.candidagenome.org/) [23], revealed that the most significant gene functions in terms of their biological process were related to transport, and the most active cellular component is the cell membrane (Table S2). This indicated that membrane transport could represent the key activated function in *C. albicans* as it grows and transforms within zebrafish during infection.

Principal component analysis (PCA) was applied to analyze the dynamic variations in these significant genes using different time points as variables. The scores of the first, second, and third PCs accounted for 79% of the observed variations in the data. The second PC accounted for an additional 12% of the variations, the third PC accounted for 5%, the fourth PC accounted for 2%, and the others accounted for less than 1%. Hence, the first two PCs could account for more than 91% of the variations in the expression dataset.

Figure 1A displays the relative contribution of different time points to the first two PCs. The first PC featured the differential expression between time points 0.5 and 1, and 12, 16 and 18 hours post infection (hpi). The second PC featured the differences between time points 2, 4, 6 and 8 h, and 0.5, 1, 12, 16 and 18 hpi. Histological analysis of the host zebrafish was performed at the exact time points corresponding with the array study. Using the host’s liver as a focus, it was found that the pathogen’s morphological features correlated well with the three different stages of fungal infection. In Figure 1C, within 1 hpi, *C. albicans* remained predominantly in yeast form and attached to the surface of the zebrafish liver. No tissue invasion was observed in the samples collected at 0.5 and 1 hpi. This period was thus defined as the adhesion phase. *C. albicans* began to form hyphae around 2 hpi, which marked the starting point for the morphological transition *C. albicans*. The fungal invasion into the liver tissue became evident from 4 to 8 hpi; therefore, the time points from 2 to 8 hpi were defined as the invasion phase. Tissue damage and extensive penetration became evident beyond 12 hpi, and the host survival rate decreased rapidly from this time point onward. This result was consistent with the observations reported in the group’s previous study [20], and thus, the time points from 12, 16, to 18 hpi were defined as the damage phase. Here PCA analysis and the histological findings were able to classified, in a phenomenological manner, experimental time points into the adhesion, invasion, and damage phases. Nevertheless, the molecular events that occurred in and between stages may be more complex and remained to be investigated.

The scores of the first two PCs accounted for more than 90% of the variations, and Figure S2A displays the scatter plot of the two scores for these significant genes. These genes were then classified into two major groups (Groups 1 and 2) by clustering principal component scores using the K-mean method and validity index. Figure 2A shows their expression profiles. There were three subgroups (Subgroups I, II, and III) with different expression patterns in Group 1. In Subgroup I, the genes were highly expressed in the adhesion phase, but their expression levels decreased when *C. albicans* entered the invasion and damage phases. The genes in Subgroup II had high positive second PC scores, and those in Subgroup III had high negative second PC scores. The expression of the genes in Subgroup II was lowest in the invasion phase, while the expression of the genes in Subgroup III was highest in the invasion phase.

There were also three subgroups (Subgroups IV, V, and VI) in Group 2. In comparison with the aforementioned data, *C. albicans* did not significantly express most of the identified genes in Subgroup IV in the adhesion phase; instead, their expression increased in the invasion and damage phases. The genes in Subgroup V demonstrated the lowest expression levels in the invasion phase, while the genes in Subgroup VI demonstrated the highest expression levels in the invasion phase.

As described previously, *C. albicans* expressed distinct groups of genes during its interactions with zebrafish. Table 1 lists the key biological processes that demonstrated an association bias toward either Group 1 or Group 2 (A complete list shown in Table S3)
with a p-value of less than 0.1. Biofilm formation, which is important in fungal pathogenesis, showed no specific bias towards group 1 or group 2 was also included as a reference. In particular, the genes related to hyphal morphogenesis, response to chemical stimuli, response to stress, organelle organization, and cell wall organization demonstrated a significant bias toward Group 1. However, the genes with a significant association bias toward Group 2 tended to function in cellular homeostasis and amino acid metabolic processes. Moreover, the response to chemical stimuli, the response to stress, organelle organization and cell walls are very common GO functions highlighted in $C.\text{ albicans}$. 

**Analysis of $C.\text{ albicans}$ genes on cell surface.** The list of significant $Candida$ genes contained 135 cell surface-related genes, which can be divided into two groups in the PC subspace. Figure 3A showed the expression profiles of these 135 genes throughout the time course of the infection. A compiled list of the genes is shown in Table S1.

There were 17 genes (subgroups II, III, V, VI in Figure 3A) that had expression profiles particularly featured in the invasion phase. Among them, previous studies have demonstrated that $ALS3$ and $INT1$ encode cell surface proteins related to $C.\text{ albicans}$ adhesion to the host cells [24,25]. $PRA1$ encodes a cell surface antigen, and disruption of the $PRA1$ gene protects $C.\text{ albicans}$ against leukocyte killing, thus increasing $C.\text{ albicans}$ virulence and organ invasion in vivo [26]. $NGT1$ encodes an N-acetylglucosamine transporter induced during hyphal growth [27], and $PGA32$ encodes a putative GPI-anchored adhesion-like protein that iron transcriptionally regulates [28]. Taken together, these findings indicated the importance of iron uptake and utilization in $C.\text{ albicans}$ virulence.

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**Figure 1. PC load bar plot and histology of zebrafish infected by $C.\text{ albicans}$.** (A) The pathogen’s load contributions to the first principal component indicated that the differences in expression between 0.5 and 1 and 12, 16 and 18 hpi represented the axes of largest variation. The load contributions to the second principal components featured the gene expression differences between time points 2, 4, 6 and 8 hpi and 0.5, 1, 12, 16 and 18 hpi. Two principal components sufficiently explained more than 93% of the variations of the zebrafish expression profiles. (B) The host’s load contributions to the first principal component indicated that the differences in expression between time points 0.5, 1, 2 and 4 hpi and 12, 16 and 18 hpi represented the axes of largest variation. The load contributions to the second principal components featured the gene expression differences between time points 6, 8 and 12 hpi and 0.5, 1, 2, 4, 16 and 18 hpi. (C) The histology of zebrafish liver indicated that $C.\text{ albicans}$ mainly grew in yeast form at 1 hpi and began to attach to the zebrafish liver. 0.5 and 1 hpi were thus defined as the adhesion phase. $C.\text{ albicans}$ began to transit to the hyphal form at 2 hpi. Invasion to the liver was evident at 4 to 8 hpi. Hence, time points 4 to 8 hpi were defined as the invasion phase, with 2 hpi defined as a transition point for morphogenesis. Beyond 12 hpi, tissue damage, more extensive penetration, and fish death occurred. Time points 12, 16 and 18 hpi were defined as the damage phase.

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As shown in Table S4, GO analysis of the biological processes linked to the cell surface-related genes revealed a number of biological processes that were related to hyphal formation and cell wall reorganization, which demonstrated a significant association bias toward Group 1. However, cellular homeostasis was found to demonstrate a significant bias toward Group 2. Seven out of the nine identified genes involved in this process were RBT5, PGA10, CSA1, SMF12, HMX1 and FTR1, all of which are known to be involved in iron homeostasis, and CRP1, which has a known function in copper homeostasis. These results suggested that iron transport and scavenging are important activities of C. albicans during the infection process.

The above analysis was summarized in Figure 4, which includes the method used, key findings and the related tables and figures. First, significant genes were first singled out by ANOVA and Bonferroni correction. GO analysis of these genes revealed that the key function is related to membrane transport. On the other hand, PCA showed that the infection can be classified into three stages according to gene expression profiles. K-mean clustering suggested that the genes profile can be cluster into two main groups: progressively expressed and progressively suppressed. Binomial tests were used to identify important GO-terms in the two main groups. For both total significant genes and cell surface-related genes, hyphal formation and cell wall reorganization demonstrated a significant association bias toward Group 1. Cellular homeostasis was found to demonstrate a significant bias toward Group 2, and most cell surface-related genes in cellular homeostasis are iron related. Therefore, we next move to identify the profiles of hyphal-related genes and iron-related genes during infection.

Hyphal-related genes of C. albicans. The analysis identified hyphal growth as the fungal pathogen’s primary process in the invasion and damage phases. There were a total of 122 hyphal-related genes. In agreement with prior research findings [6,8,10]...
and CGD annotation, most of these genes have been reported to contribute to \textit{C. albicans} virulence in animal models other than zebrafish, and have thus been identified as essential virulent genes for \textit{C. albicans}. These genes could also be divided into two groups in the PC subspace (Table S1). Figure 3B shows their expression profiles throughout the time course of the infection.

Group 1 contained 86 genes, and Group 2 had 36 genes. There were six genes that had profiles featured particularly in the invasion phase. Among them, it was found that \textit{C. albicans} transiently expresses \textit{SHA3}, \textit{FCR1} and \textit{DFG16} in the invasion phase. In terms of gene function, \textit{SHA3} has a known involvement in glucose transport [29], while \textit{FCR1} encodes a putative zinc cluster transcription factor [29]. \textit{DFG16} is required for host tissue invasion [30]. Conversely, \textit{C. albicans} transiently expresses \textit{CDC14}, \textit{CLB3}, and \textit{BUB2} in the invasion phase. In terms of gene function, \textit{CDC14} is related to cell cycle function, \textit{CLB3} is required for mitotic exit [31], and \textit{BUB2} participates in the mitotic cell cycle spindle orientation checkpoint [31].

Multiple signaling and regulatory pathways control \textit{C. albicans} yeast-to-hyphal transition. This study identified 122 significant filamentous related genes. Previous studies have mapped many of these genes onto pathways known to control morphological transition, and these genes expressed significantly during infection are arranged in Table 2. \textit{GST20} and \textit{CEK1} encode components of the mitogen-activated protein kinase (MAPK) signaling pathway [10]. Moreover, \textit{CDC42}, \textit{CDC24}, and \textit{BEM3} gene products function upstream of the MAPK pathway [32] and are expressed during the adhesion phase. \textit{HOG1}, a MAP kinase involved in osmotic, heavy metal, and core stress responses [33,34], is also activated during the adhesion phase. \textit{SSK1} and \textit{CHK1}, which are response regulators of the two-component system, are activated during the damage phase [33,36].

\textit{C. albicans} expressed the downstream target of the protein kinase C (PKC) pathway, \textit{MKC1}, which is involved in contact-induced invasive filamentation, during the adhesion phase. Of the remaining essential biofilm formation genes, \textit{C. albicans} up-regulated \textit{RBT5} and transiently expressed \textit{ALS3} during the invasion phase. Of the genes involved in the pH response signaling pathway, \textit{C. albicans} was found to express \textit{RIM21} during the damage phase and \textit{PHR1} during the adhesion phase. \textit{Rim21} may function as a sensor of alkaline pH [37], and \textit{Phr1} is a putative cell surface glycosidase with involvement in cross-linking of the beta-1,3- and beta-1,6-glucans [38].

The results of this study further revealed the modulation of several components in the \textit{C. albicans} TOR pathway. This was significant, as the target of the rapamycin (TOR) signaling pathway regulates a wide variety of cellular processes in response to nutrients. Of these, \textit{C. albicans} expressed \textit{GLN3} during the damage phase, and expressed \textit{RHB1}, \textit{GAP1}, \textit{MEP2} and \textit{SIT4} during the adhesion phase.

\textit{C. albicans} activated other hypha-related genes, such as \textit{SIN3} and \textit{SET3}, during the adhesion phase. \textit{SIN3} is known to encode a key component of a specific histone deacetylase complex. It can down-regulate true hyphae formation via binding to the \textit{EFG1} promoter [39]. \textit{Efg1} is a transcriptional regulator and activates filamentous growth in \textit{C. albicans}. On the other hand, the gene product of \textit{SET3} forms a histone deacetylase complex that is involved in regulating \textit{C. albicans} morphogenesis and virulence [40].

The present study’s analyses of \textit{C. albicans} also identified genes related to calcium. It was found that \textit{C. albicans} would express calmodulin (\textit{CMD1}) and the catalytic subunit of calcineurin (\textit{CMP1}). \textit{C. albicans} also up-regulated the expression of \textit{MID1}, which encodes a putative calcium channel of the high affinity calcium uptake system, during the damage phase. \textit{MID1} plays a role in thigmotropism (contact sensing) and the directional hyphae tip response of \textit{C. albicans} [41].

Taken together, the results indicated that \textit{C. albicans} exhibits dynamic gene regulatory controls in many well-known signaling pathways to coordinate various functions during the adhesion phase, but that it will then progressively suppress them. This result was consistent with the finding of yeast-to-hyphal transition occurring in the early stage of infection (Figure 1C).

\textbf{Iron-related genes of \textit{C. albicans}.} The analysis of cell surface-related genes identified iron transport and scavenging as the primary processes. The expression of these iron-related genes mainly occurred during the invasion and damage phases. Using the

\textbf{Table 1.} \textit{C. albicans} genes cluster by PCA subspace.

\begin{table}
\centering
\begin{tabular}{|l|c|c|c|c|}
\hline
\textbf{GO term} & \textbf{Group 1} & & \textbf{Group 2} & \\
 & \textbf{No. of Genes} & \textbf{p-value} & \textbf{No. of Genes} & \textbf{p-value} \\
\hline
DNA metabolic process & 53 & 0.000 & 2 & 1.000 \\
vesicle-mediated transport & 82 & 0.000 & 16 & 1.000 \\
response to stress & 82 & 0.000 & 33 & 0.999 \\
filamentous growth & 86 & 0.000 & 36 & 0.999 \\
transport & 152 & 0.000 & 76 & 0.999 \\
hyphal growth & 44 & 0.001 & 14 & 0.999 \\
response to drug & 55 & 0.001 & 21 & 0.997 \\
response to chemical stimulus & 86 & 0.003 & 41 & 0.995 \\
translation & 15 & 0.003 & 3 & 0.985 \\
ribosome biogenesis & 11 & 0.006 & 2 & 0.968 \\
protein catabolic process & 37 & 0.015 & 16 & 0.970 \\
cellular amino acid metabolic process & 19 & 0.943 & 25 & 0.030 \\
cellular homeostasis & 8 & 0.987 & 17 & 0.004 \\
\hline
\end{tabular}
\caption{\textit{C. albicans} genes cluster by PCA subspace.}
\end{table}

This table lists the key biological processes that demonstrated an association bias toward either Group 1 or Group 2 with a \textit{p}-value of less than 0.05 (A complete list was shown in Table S3). doi:10.1371/journal.pone.0072483.t001
The literature search approach described in Material and Methods under the specific functional gene list section, the iron-related genes list was found to contain 238 genes, 61 of which had significant expressions. Of these, 25 were in the Group 1 cluster, while 36 were in the Group 2 cluster within the PC subspace (Table S1). Figure 3C provides a list of the genes in these two groups.

Figure 3. Expression profiles of the cell surface-related, hyphal-related, and iron-related genes in *C. albicans*. The list of significant genes contained 135 cell surface-related genes. PCA divided these into two main groups. (A) shows the cell surface-related gene expression profile. 16 genes, COX11, ERG11, orf19.781, ALS3, CNT, orf19.3983, orf19.6435, SMF12, DFG16, orf19.4947, SCO1, PRA1, PLB5, INT1, NGT1 and PGA32 were particularly featured in the invasion phase. The list of significant genes also contained 122 hyphal-related genes. PCA divided these into two main groups. (B) shows the hyphal-related gene expression profile. Six genes were particularly featured in the invasion phase. Of these, *C. albicans* transiently suppressed SHA3, FCR1 and DFG16 during the invasion phase. Analysis of the cell surface-related genes identified cellular hemostasis as a primary process on the cell surface, and most of these genes were iron homeostasis-related. The list of significant genes contained 61 iron-related genes. PCA divided these into two main groups. (C) shows the iron-related gene expression profile. Nine genes were particularly featured in the invasion phase. *C. albicans* transiently suppressed HEM14, SGE1, YHB1, DFG16 and SEO1, and transiently expressed ALS3, ERG11, INT1 and PGA32 in the invasion phase.

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C. albicans is known to regulate different genes to utilize and uptake iron by sensing the iron concentration in its environment. Accumulated evidence to link these genes' induction to the surrounding iron concentration has been described in previous studies [28,42–49]. Genes related to iron uptake and utilization that were expressed significantly during infection [18,28,42,45–58] are arranged in Table 3 according to whether they are expressed in high/low ex vivo experiments, and whether they belong to group 1 or 2 in in vivo experiments. There is no correlation between two classifications. This shows the importance of understanding the in vivo response of iron-related genes during infection.

We also mapped present significant genes into iron-related pathway to identify the key factor during infection. There are at least three known strategies for C. albicans to acquire iron from its host: the siderophore uptake system, the reductive system, and the heme-iron uptake system. The primary iron source appears to be the iron-storage protein, ferritin [42]. Many of these genes have been mapped onto these pathways controlling morphological

![Diagram](image)

**Figure 4. C. albicans genes analysis procedure diagram.** The analysis of C. albicans was summarized in this figure, which includes the method used, key findings and the related Tables and Figures. First significant genes were first singled out by ANOVA and Bonferroni correction. GO analysis of these genes revealed that the key function is related to membrane transport. PCA showed that the infection can be classified into three stages according to gene expression profiles. K-mean clustering suggested that the genes profile can be cluster into two main groups: progressively expressed and progressively suppressed. Binomial tests were used to identify important GO-terms in the two main groups. For both total significant genes and cell surface-related genes, hyphal formation and cell wall reorganization demonstrated a significant association bias toward Group 1. Cellular homeostasis was found to demonstrate a significant bias toward Group 2, and most cell surface-related genes in cellular homeostasis are iron related.

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**Table 2.** Significant C. albicans genes in signaling pathway of yeast-to-hyphae transition.

| Path | Group 1 | Group 2 |
|------|---------|---------|
|      | Progressively suppressed | Transient | Progressively expressed |
| MAPK | CST20, CEK1, CDC42, CDC24, BEM3, HOI1 | | SSK1, CHK1 |
| PKC  | MKC1 | ALS3 | RBT5 |
| pH   | PRH1 | | RIM21 |
| TOR  | RBH1, GAP1, MEP2, SIT4 | | GLN3 |

Map significant filamentous related genes onto well-known controlling morphological transition pathways.

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transition. Als3, which is known to play an important role in hyphae formation and invasion, is needed to acquire iron from the host’s ferritin. Based on the dynamic gene expression data, it was found that C. albicans only transiently expressed ALS3 in the early invasion phase, while FTR1 and other ferric reductases (FRP1, FRP2, and GRE2) were progressively induced during the invasion phase. Their expressions were then sustained in the damage phase. These data indicated that the uptake of iron had already started in the early invasion phase. On the other hand, C. albicans expressed HMX1 in the damage phase, and up-regulated PGA10 and RBT5 during the invasion and damage phases. This indicated the possible uptake of iron from the host’s hemoglobin as the zebrafish suffered massive hemorrhaging during the damage phase. The microarray profiles of ALS3, FTR1 (reductive system), RBT5, and HMX1 (heme-iron uptake system) are shown in Figure 5. This summarized the finding that the reductive uptake was turned on during the adhesion and invasion stages, while the heme-iron uptake become important in the damage phase.

**Table 3. Profile of iron-related genes during infection and reference.**

| Reference iron environment | Progressively suppressed | Transiently suppressed | Progressively expressed | Transiently expressed |
|---------------------------|--------------------------|------------------------|------------------------|-----------------------|
| High                      | KAR2, BGL2, RFG1, SEC12, orf19.2452 | HEM14, YHB1 | MDH1, PGA10, CHT2, ISU1, orf19.2048, BIO2, orf19.413, FUM11, SBA1, SPL1, HSP90 | ERG11, PGA32 |
| Low                       | SAP10, GEA2, GCS1, MET3, orf19.1917, orf19.2244, orf19.3216 | SGE1, SEO1 | HMX1, RBT5, FTR1, SOD5, CSA1, FRP1, FTH1, CHA1, GRE2, GCV1, IHD2, SAP99, orf19.6530 | INT1 |
| Unknown                   | FNE7, CS11, MIG1, MAC1, HOG1, SHE3, RLJ1, SOD3, HEM15, orf19.5369 | DFG16 | VPS28, MYOS, FRP2, IRD1, SOD1, SOD2 | ALS3 |

Compare profile of iron-related genes with those previous researches, which link these genes’ induction to the surrounding iron concentration.

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Dynamic gene expression analysis of zebrafish host response during *C. albicans* infection

The present study applied the same statistical methods to analyze the zebrafish host’s dynamic gene expression profile during *C. albicans* infection. The expression profile contained a total of 43,803 probes. Similarly, one-way ANOVA was applied to detect significant gene expression variations across the nine-point time course. Figure S1B illustrates the numbers of significant genes identified at different *p*-values. There were a total of 10,826 significant probes with a *p*-value less than 0.01. Gene annotation was carried out using the Database of Visualization and Integrated Discovery (DAVID) v6.7 [59] and ZFIN [60]. It shall be noted that, unlike the *Candida* pathogen database, there was less bioinformation available for zebrafish. The resulting list of significant genes contained 6,150 genes. Many known genes associated with host defense or immune response mechanisms were in this list. However, the following discussion focused on a list of 576 significant genes (Table S5) that fulfilled the stringent criterion of a *p*-value less than 0.01/43803 (Bonferroni correction).

PCA was applied to analyze the dynamic variations in these significant genes using different time points as variables. The scores of the first PC accounted for 84% of the observed variations in the data, while the second PC explained an additional 9% of the variations. The third PC explained 2%, and the others explained less than 1%. Therefore, the first two PCs could account for more than 93% of the variations in the expression dataset.

Figure 1B displays the contribution plots of these two principal components. The first PC featured the differences in gene expression between time points 0.5, 1, 2 and 4 hpi, and 12, 16 and 18 hpi. The second PC featured the expression differences between time points 4, 6, 8 and 12 hpi, and 0.5, 1, 2, 16 and 18 hpi. This result indicated that the invasion phase in zebrafish was slightly delayed compared to that in *C. albicans*.

Figure S2B illustrates the loci of significant zebrafish genes in the PC subspace. These could be clustered into two main groups according to the validity index. Figure 2B shows the expression profiles of the two groups. The genes in Group 1 (334 probes, 267 genes) demonstrated high expression in the adhesion phase, but this expression started to decline upon entering the invasion and damage phases. The genes in Group 2 (349 probes, 309 genes) demonstrated low expression in the adhesion phase, but this expression increased in the invasion and damage phases.

Gene ontology analysis of these 576 significant zebrafish host genes was performed using DAVID. The analysis revealed that the cluster of genes with the highest enrichment score was related to iron binding (Table S6). The cluster with the second highest enrichment score was related to hemostasis. The high enrichment score relating to hemostasis was supported by observations of frequent internal bleeding during the experiments.

**Figure 5. Expression profiles of significant *C. albicans* genes on well-known iron uptake pathway.** In reductive system, ALS3, needed to acquire iron from the host's ferritin, transiently expressed in the early invasion phase. FTR1, a high-affinity iron permease, was progressively induced during the invasion phase, and expression was then sustained in the damage phase. In heme-iron uptake system, C. albicans up-regulated RBT5 during the invasion and damage phases and expressed HMX1 in the damage phase. These data indicated that the uptake of iron had already started in the early invasion phase and the possible uptake of iron from the host’s hemoglobin as the zebrafish suffered massive hemorrhaging during the damage phase.

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Iron-related genes of zebrafish. The previously mentioned analyses of the \textit{C. albicans} gene expression profiles identified iron transport and scavenging as important processes during infection. Interestingly, GO analysis of the significant genes in the host zebrafish also indicated that iron homeostasis was important biological process too. Of the 576 significant zebrafish genes, 24 displayed GO associated with iron binding and iron transport.

Figure 6A shows the expression profiles of the iron-related genes throughout the infection time course. Group 1 contained 18 genes that the zebrafish progressively suppressed, and Group 2 contained six genes that were progressively expressed, including.

It was worth mentioning that the zebrafish progressively suppressed most of the iron-related genes during infection. It should also be noted that most of the iron-related genes were categorized based on informatic annotations instead of experimental data. Nonetheless, one previous wet bench study has identified SLC25A37 to be related to iron transporter activity and hemeprotein synthesis [61].

Immune-related genes of zebrafish. The immune response is a key process for zebrafish to defend themselves against \textit{C. albicans} infection. Of the 576 significant genes identified, 13 had GO related to immune response and immune system development.

Figure 6B shows the expression profiles of these genes throughout the time course of the infection. Group 1 contained three genes. Previous studies have demonstrated that HBL4 and VTNB encode components of the extracellular matrix [62–64]. HBL4 and VTNB are involved in damaged cell repair and are significantly expressed during the adhesion phase, but become progressively suppressed in the damage phases. Group 2 contained the other 10 genes. Among them, \textit{IL1B}, \textit{TNFA} and \textit{TNFB} are well-known marker genes of the immune response. Further analysis also identified the expression of the growth factor gene FGF21, which implied that zebrafish may activate a healing mechanism to repair the damage caused by \textit{C. albicans} infection. This was supported by the observed up-regulation of RUNX1, which may promote the production of blood cells following tissue penetration by \textit{C. albicans}. It was noteworthy that the observed immune response occurred immediately after the initial adhesion phase of the infection.

In contrast to previous studies that used cell lines, tissues and organs to pass as a host system for microbial infection, this study utilized a whole organism with complete immune capability. The GO analysis revealed that only a small set of known immune-related genes were significantly expressed the zebrafish. This indicated that the immune response was only one of numerous simultaneous biological activities vigorously regulated in the host during infection. The complex interaction between the immune system and the other physiological processes in the host warrants further investigation in the future.

The above analysis was summarized in Figure 7, which includes the method used, key findings and the related Tables and Figures. First, the significant genes were singled out by ANOVA and Bonferroni correction. GO analysis of these genes revealed that the key function is related to iron-related function. PCA and K-mean clustering suggested that the genes profile can be cluster into two main groups: progressively expressed and progressively suppressed. Finally, binomial tests were used to identify the bias of specific functions. Iron-related genes demonstrated a significant...
bias toward Group 1 p-value 0.0011. Immune-related genes demonstrated a significant association bias toward Group 2 with p-value 0.0215. These results indicated that the onset of immune in the host is highly correlated with failure of iron regulation.

Comparison between Candida gene expression profiles in different hosts

This present study also compared Candida genome expression profiles in other host models to investigate possible virulence genes. There are several published studies providing accessible transcription profiles of C. albicans during infection in different living hosts, such as mice and rabbits [17–19,65]. This study chose the C. albicans genome-wide data set from Thewes et al. (2007) [18], whose microarray dataset has a complete Candida gene list and a 4 time point (0, 0.5, 3, 5hpi) series profile generated from samples collected from the livers of mice infected with Candida. To ensure consistency, this study applied the proposed method to re-analyze Thewes’s gene expression dataset and identify two groups of pathogen genes, including 288 up-regulated genes and 185 down-regulated genes. Figure 8A shows that the total intersection of the conserved genes from this work and Thewes et al. consisted of 130 genes. Figure 8B and Figure 8C show 39 up-regulated genes and 45 down-regulated genes had the same expression profile in both the zebrafish and the mouse models. The entire list is shown in Table S7.

Among the 84 genes with common dynamic expression profiles, the functions of the genes with the top 15 fold change were studied. In this list, SAP5 is associated with the utilization of protein as nitrogen source and virulence [66]. RHR2, regulated by HOG1 and NRG1, is a response to biofilm and hyphal growth [67]. SNZ1, CHT2, ZRT1, and PCK1 are associated with hyphal growth [68–70]. SIM1, described as a macrophage-down-regulated gene, was repressed both in mice and fish. In terms of the iron-related genes, SOD5, PGA7, FTR1, and RBT5 were identified. FTR1 is a ferric reductase, RBT5 is known as a hemoglobin receptor, and PGA7 is described as part of a hemoglobin-receptor gene family [71]. orf19.6084, orf19.449, and orf19.5848 are described as biofilm-induced genes [72]. There is no description for orf19.215, but it is an ortholog to ScAVO2, which is related to Tor2p kinase and other proteins.

Interestingly, there were 29 genes out of 1307 genes related to temperature in the fish model, and 14 genes out of 473 genes in mouse model. There were only five temperature-related genes found in both models (MYO2, SRV2, PKG1, SMT3 and MKC1). A close examination of their expression profiles revealed that only MYO2 was progressively suppressed in both models, and that C. albicans up-regulated the other four genes in mice and down-regulated them in fish. This may have been caused by the body temperature difference between mice (36.9 degree Celsius) and zebrafish (28 degree Celsius). In a previous study, MYO2 was found to be not only related to temperature but also to cytoskeletal polarity, migration, and hyphal growth, therefore MYO2 may not be directly affected by temperature [73,74].

To summarize, C. albicans will turned on a set of the specific virulence genes, which includes genes related to pathogenesis and iron homeostasis, in both fish and mouse models. These genes not
only activated the hyphal growth, but also help the pathogen to uptake iron in the host's microenvironment. It should be pointed out that our dataset include the whole body host response to pathogen, but the dataset of Thewes et al. (2007) was an organ-specific response. Hence there must be caution against too much emphasis on the similarity between the datasets. However, these Candida genes with common expression profiles in different host species may indeed be truly important during C. albicans infection.

Conclusions

This study analyzed a set of time-lapse genome expression data between the Candida pathogen and zebrafish that captured snapshots of dynamic molecular events during infection. It was found that C. albicans turned on yeast-to-hyphae morphogenesis to invade the host, and that many cell surface-related and hyphal-related genes were activated during the early stages of infection.

As expected, the zebrafish progressively induced their immune systems to defend against the aggressor. One of the host’s immune defenses was to restrict the iron available to the pathogens [75–77]. Many of the iron-related genes were suppressed in the late phase. A contrary example, SLC25A37, which is responsible for pumping iron out of the cell for hemooprotein synthesis [61], was consistent with the hypothesis that the zebrafish host was trying to sequester iron at the initial phase of infection, and that its late expression in the damage phase was caused by massive hemorrhaging. Also, the zebrafish expressed immune related genes much more significantly during the late stage of infection. In response to this defense mechanism, the reductive and heme-iron uptake were found to be important strategies for C. albicans to acquire iron from the host [42]. The genes responsible were progressively induced, especially after the collapse of the iron restricting mechanism in the fish host. The expressions of FTR1, a reductase in the reductive system, and RBT5, a hemoglobin receptor in the heme-iron system, were found to be similar in both the fish and mouse models. The microarray profiles were validated using real-time PCR. The expression patterns of the RBT5, CHT2, MNT1 and PMT1 C. albicans genes (Figure 9A) and the TNFA, SLC25A37, VT/NB and CIP4V2 zebrafish genes (Figure 9B) were consistent with those found in the DNA microarray analyses.

Kuo et al. 2013 [22] study the host-pathogen interaction dynamics, based on available data-base information of PPI network. A time series modeling method and Aidaike Information Criterion were used to determine the active inter-species and intra-species networks in adhesion stage and invasion stage, respectively. It shall be noted that the division of the two stages was phenomenologically based. The top ranked proteins found for C. albicans are mainly related to hyphal growth. The top ten proteins with active interactions in the host are mainly related to apoptosis and innate immunity. In contrast, the present study used the gene expression profiles to support statistically the phenomenological classification of the fungal infection processes into distinct phases (adhesion, invasion, and damage) throughout the pathogen's infection episode. Furthermore, we suggest iron competition is important for the emergence and progression of pathogen virulence. The pathogen invasion process is accompanied by the failing of iron homeostasis function in the host and the switching of iron uptake pathway of the pathogen. Although the interaction between the immune system and other physiological processes of zebrafish require further investigation in the future, this present work demonstrated the importance of correlation of intra-species regulatory activities from stage to stage. Such knowledge may facilitate the development of novel approaches to attenuate C. albicans virulence.

Materials and Methods

Candida albicans and zebrafish

Adult AB strain zebrafish were used for the experiments. Their maintenance and care after infection were performed according to the procedures described previously [20]. All of the zebrafish use and experimental protocols in this research were reviewed and approved by the Institutional Animal Care and Use Committee of National Tsing Hua University (IRB Approval NO. 098086). C. albicans SC5314 strain was used for injection and was prepared as described previously [20].

Infection and histological analysis

For the microarray analysis, each fish was intraperitoneally injected with $1 \times 10^6$ C. albicans cells suspended in 10 μl sterile 1X phosphate-buffered saline (PBS) after being anesthetized using 0.17 g/ml of tricaine (Sigma). At each time point (0.5, 1, 2, 4, 6, 8, 12, 16 and 18 hpi), three infected fish were collected, sacrificed by immersion in ice water, frozen in liquid nitrogen, and then maintained at −80°C. Although using whole infected fish to obtain transcriptional profiles could result in the loss of specific information, such as tissue and organ data, it allowed the investigation of the entire genome response during C. albicans infection. For the survival assay, the total injected yeast form cell number was $10^9$ cells. For histological analysis, fish were collected at designated time points, and then fixed, sectioned, stained and observed according to the methods used previously [20].

C. albicans and zebrafish RNA purification

The zebrafish infected with Candida albicans were treated with Trizol® reagent (Invitrogen, USA), ground in liquid nitrogen using a small mortar and pestle, and then disrupted using a MagNA Lyser system (Roche) that used glass beads (cat. no. G6772-100G,
Sigma) shaken at 5,000 rpm for 15 sec. After phase separation was performed by adding chloroform, the total RNA was purified using an RNeasy Mini Kit (Qiagen, Germany). Purified RNA was quantified at OD260nm using an ND-1000 spectrophotometer (Nanodrop Technology, USA) and qualitized using a Bioanalyzer 2100 (Agilent Technology, USA) with an RNA 6000 nano labchip kit (Agilent Technologies, USA).

Figure 9. Validation of expression patterns by real-time polymerase chain reaction. (A) The C. albicans gene expression patterns (RBT5, CHT2, MNT1, and PMT1) revealed by the DNA microarray are shown on the top. Real-time PCR of C. albicans genes (RNA isolated from three infected zebrafish per group) are shown in the bottom. C. albicans expressed RBT5 to use iron in the invasion and damage phase (after 6 hpi), expressed CHT2 in the late stage (after 12 hpi), and activated MNT1 and PMT1 in the adhesion phase (0.5 and 2 hpi). (B) The zebrafish gene expression patterns (TNFA, SLC25A37, VTNB, and CYP4V2) revealed by the DNA microarray are shown on the top. Real-time PCR of the zebrafish genes (RNA isolated from three infected zebrafish per group) are shown on the bottom. The zebrafish progressively expressed TNFA and SLC25A37 during the infection and turned on VTNB and CYP4V2 in the adhesion phase (0.5, 1 hpi). The profiles of the real time PCR were identical and could validate the present genome microarray in both the host and the pathogen. The asterisks above the bar indicate that the level of target gene expression was significantly different from the levels of expression in the other time points: * = P<0.2; ** = P<0.05; *** = P<0.01.

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Preparation of _C. albicans_ custom array and zebrafish array

For the _C. albicans_ gene expression analysis, probes were designed using eArray of Agilent technologies. Along the design process, 6,205 _Candida albicans_ transcripts (Assembly 21, from the _Candida_ genome database) were uploaded to eArray and designed using base composition methodology. Duplicated target sequences were removed and the resulting 6,202 probes were designed. The custom microarray was manufactured in an 8×15k format by in situ synthesis of the oligonucleotide probes (Agilent Technology). Each array consisted of 6,202 _Candida albicans_ specific probes, which were printed in duplicate. The _C. albicans_ gene expression analysis used Agilent zebrafish V2 oligo arrays (Agilent Technology), which were manufactured in a 4×44K format. Control experiments were performed to that ensure the _C. albicans_ samples would not cross-hybridize to the zebrafish microarrays and that the _C. albicans_ samples would not cross-hybridize to the _C. albicans_ microarrays. In this experiment, the cross-hybridization between different samples and microarrays was not significant (data not shown).

Microarray experiments

A total of 1 µg of the total RNA was amplified using a QuickAmp labeling kit (Agilent Technologies, USA) and labeled with Cy3 (CyDye, PerkinElmer, USA) during the _in vitro_ transcription process. A total of 0.625 µg of Cy3 cRNA for the _C. albicans_ array and 1.65 µg of Cy3 cRNA for the zebrafish array was fragmented to an average size of approximately 50 to 100 nucleotides through incubation in a fragmentation buffer at 60°C for 30 minutes. The fragmented labeled cRNA was then hybridized to an oligo microarray at 60°C for 17 h. After washing and drying using a nitrogen gun, the microarrays were scanned using an Agilent microarray scanner (Agilent Technologies, USA) at 535 nm for Cy3. The scanned images were analyzed using Feature extraction 9.5.3 software (Agilent Technologies, USA), and image analysis and normalization software was employed to quantify the signal and background intensities for each feature. The raw data were uploaded onto the NCBI Gene Expression Omnibus Database (GSE32119).

Microarray data analysis

The set of zebrafish arrays were processed separately from the _C. albicans_ arrays. The Loess normalization curve for the _C. albicans_ data was fitted using every probe on the array, while the Loess curve for the zebrafish data was fitted using the 485 control probes. After the pre-processing step with Loess normalization, the mean intensity levels for each of the 6,202 pairs of duplicated probes on the _C. albicans_ array were calculated as representatives. For each of the probes, one-way ANOVA was adopted to screen for significant time variations across the nine time points in either set of data. The significance level was adjusted using the Bonferroni correction, which is used for data sets with extremely large number of samples, to achieve a family-wise error rate of 0.01. The adjustment resulted in 1,307 significant probes (out of the total 6,202) for the _C. albicans_ data (Table S1) and 683 probes (out of the total 43,803) for the zebrafish data (Table S5).

Principal component analysis

Principal component analysis (PCA) uses an orthogonal transformation to convert a set of sample observations of potentially correlated variables into a set of values of uncorrelated variables called principal components [78]. Let microarray as

\[ \{X\}_{t\times J} \]

be a set of _J_-dimension multivariate data with sample points \( I \) \[ \{X\}_{t\times M} = \{I\}_{t\times M} \{P\}_{M\times J} + \{e\}_{t\times J} \].

It can be compressed into a set of _M_-dimension \((M<J)\) reduced-order data \( \{I\}_{t\times M} \) known as scores by a transformation defined by the matrix \( \{P\}_{M\times J} \), and \( e \) is noise. There were nine PCs in _p_ that could explain the microarray variations. In this study PCA was used to analyze the variations of significant genes by treating different time points as variables and different genes as samples. The relative contributions of the the _m^th_ PC can be computed as:

\[
\text{relative contribution} = \frac{\text{tr}(\{u\}_{m\times m})}{\text{tr}(\{X\}_{m\times m})} \times 100%
\]

The principle component analysis first identified that load vector with the largest contribution, and other load vectors with decreasing contributions in a sequential manner. Using this method, the key dynamic features of the data set can be captured.

K-means clustering and validity index

The distributions of genes in the _M_-dimension PC subspace can be clustered into different groups by standard K-mean clustering analysis of the score \( \{I\}_{t\times M} \). In K-mean analysis the number of clusters must be specified a priori. Let \( z_j, k = 1...K \) be the centers of K clusters \( \{C_1,...,C_k\} \) determined in the K-mean analysis. The validity ratio is defined as the ratio of average intra-group distance, i.e. the average distance between sample points and their cluster centers, and intergroup distance, i.e. the minimum distance between two cluster centers:

\[
\psi(K) \equiv \frac{1}{J} \sum_{j=1}^{J} \sum_{k \in C_k} \|z_j - z_k\| \quad \min_{k,k' \in C_k} \left( \frac{\|z_k - z_{k'}\|}{\|z_k - z_l\|} \right)^2
\]

\[
\text{inter} = \min_{k<k'} (\|z_k - z_{k'}\|)^2, k, k'=1,2,...,K
\]

The optimum number of cluster \( K^* \) is determined by minimizing the validity index \( K^* = \arg\max_k \psi(K) \).

Binomial test for group gene ontology annotation

We have determined that the microarray data can categorized into two major groups in terms of dynamic response. The binomial tests were used to determine whether a process was substantially biased toward a certain group. Let \( g \) represent the probability of finding a gene in Group 1. For example, there were 1,307 significant genes in _C. albicans_, of which 734 were in Group 1 and 573 were in Group 2; hence, \( g = 734/1307 \). Let \( n \) represent the total number of genes related to a specific biological process, e.g. \( n = 55 \) for the DNA metabolic process as shown in Table 1. Let \( m \) represent the number of genes related to this process found in Group 1, \( q = n-m \) for or the DNA metabolic process as shown in Table 1. The _p_-value of bias toward Group 1 was defined as the cumulative probability of _m_ or fewer genes being distributed into Group 1, when _n_ genes are randomly distributed into Group 1 and Group 2 according to the probabilities _g_ and _1–g_.
For the example of DNA metabolic process, the p-value is 0.000. The smaller the p-value, the smaller is the probability of observing the given distribution by chance. Hence the bias of DNA metabolic process towards group 1 is definite, as well as the bias of cellular homeostasis towards group 2 (a p-value of 0.004 in Table 1). On the other hand, data in Table 1 showed that we cannot determine whether genes related to biofilm process is bias toward group 1 (p-value 0.430) or group 2 (p-value 0.333). For regular statistical testing with small number of samples, a p-value of 0.05 is used. This p-value was thus used in the binomial test for identification of bias distributions of a small set of genes among two different groups.

Specific functional gene list
To collect all possible receptors and sensors on the C. albicans membrane, text-mining in the GCD was performed, using the keywords “plasma membrane”, “cell surface”, and “cell wall”, in order to research the GO terms “transport” and “membrane” from Table S1. The specific keywords identified a total of 533 genes, which were denoted as cell surface-related genes.

Hyphal-related biological processes were also researched by text-mining using the keywords “filamentous growth”, “hyphal growth”, and “pseudohyphal growth”. This identified 431 genes, which were named filamentous related genes.

As shown in Table S4, the significant biological processes demonstrating the bias toward Group 2, progressive expression, was cellular homeostasis. Conducting searches for possible previously identified iron-induced genes, iron phenotypes genes, predicted iron-related genes [18,46,50–55], iron-responsive regulators [56–58], iron uptake genes [42], iron utilization genes [45,47–49] and iron-modulated genes [28] obtained 238 genes, which were named iron-related genes.

RNA isolation and quantitative real-time PCR
The control fish and the infected fish (infected with 10 μl 1×10⁸ CFU/ml C. albicans cells) were sacrificed by immersion in ice water at different time points and homogenized in liquid nitrogen. The total RNA was extracted using TRIzol reagent according to the manufacturer’s instructions (Invitrogen), and the RNA quality was analyzed by gel electrophoresis. For cDNA synthesis, 6.25 μg of the total RNA was pretreated with DNase I (Invitrogen) and then reverse transcribed using the SuperScript III enzyme (Invitrogen) in a 50 μl reaction mixture. Quantitative real-time PCR was carried out using the 7500 real-time PCR system (Applied Biosystems). The primers used in this study are listed in Table S8. Briefly, each 15 μl reaction mixture contained 25 ng cDNA, 7.5 μl SYBR green PCR master mixture (Applied Biosystems), 0.2 μM forward primer, and 0.2 μM reverse primer. The reactions were performed using 1 cycle of 95°C for 10 min followed by 40 cycles of 95°C for 15 s and 60°C for 1 min.

Supporting Information
**Figure S1** C. albicans and zebrafish genes at different p-values. (A) Venn diagram of the significant genes in C. albicans. There were a total of 4,827 significant genes with a p-value of less than 0.01, and there were 4,827 significant genes with a p-value of less than 0.1/6202 (Bonferroni correction). This study focused on 1,307 genes that fulfilled the stringent criterion of a p-value less than 0.01/6202. This list included several genes related to C. albicans hyphal formation, such as MIG1, RFG1, ECE1, and CEK1. (B) Venn diagram of the significant genes in zebrafish. There were a total of 36,292 significant genes with a p-value of less than 0.01, and there were 6,150 significant genes with a p-value of less than 0.1/43803. This study focused on 683 genes that fulfilled the same criterion used in the C. albicans analyses; namely, a p-value less than 0.01/43803 (Bonferroni correction). This gene list included key genes related to the immune response, such as IL1B, TNFA, and TNFB.

**Table S1** C. albicans significant genes list. Applying a stringent criterion of a p-value less than 0.01/6202 = 1.61×10⁻⁶ (with Bonferroni correction) further reduced the target gene pool and identified a total of 1,307 significant genes. The gene list also shows classifications according to groups in the PC subspace and cell-surface, iron and hyphal related functions.

**Table S2** C. albicans gene ontology analysis. Gene ontology annotation of 1,307 genes performed using the Candida Genome Database and the GO Slim Mapper terms “biological process” and “cellular components”. The most significant biology process GO set related to transport. The most significant cellular component GO set related to the cell membrane.

**Table S3** GO comparison of total C. albicans significant genes group. The table shows the GO of identified biological processes and demonstrating a significant bias in Group 1 or Group 2. Binomial distribution was used to calculate the p-value, assuming that significant genes of a specific process were randomly distributed between two groups.

**Table S4** GO comparison of C. albicans cell surface related genes group. The table shows the GO of identified biological processes of C. albicans cell surface related genes and demonstrating a significant bias in Group 1 or Group 2.

**Table S5** Zebrafish significant genes list. A list of 576 significant genes that fulfilled the stringent criterion of a p-value less than 0.01/43803 (Bonferroni correction); the same criterion used in C. albicans data analyses. The compiled list also shows classifications according to groups in the PC subspace.

**Table S6** Zebrafish gene ontology analysis. Gene ontology analysis result of 576 significant zebrafish host genes using DAVID.

**Table S7** Identical expression profile of C. albicans genes in mouse host. The proposed method was used to re-analyze the gene expression profiles in the C. albicans genome wide data set in Thewes et al. (2007) [18] and cluster them into two
group of 288 up-regulated genes and 185 down-regulated genes. Among them, 39 up-regulated genes and 45 down-regulated genes had the same expression profile in the zebrafish model. The 84 genes are shown in this table.

**Table S8** Sequences of primers used for real-time quantitative PCR. The primers used in this study are listed in this table.

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**Author Contributions**

Conceived and designed the experiments: CCC FCH HFC. Analyzed the data: YYC SCP DSHW WPH. Contributed reagents/materials/analysis tools: CCC FCH HFC YJC. Wrote the paper: YYC YJC CYLDSHW.

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