Bioinformatic analysis of the pathogenic mechanism of talaromyces marneffei infection

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Abstract

**Background:** Talaromyces marneffei (T. marneffei), known as a significant pathogen in patients with AIDS in Southeast Asia, is a dimorphic fungus, which can cause deadly systematic infection in immunocompromised hosts. What is more, the dimorphic phase transition has been reported as a conspicuous process linked with virulence. Interestingly, the yeast form was found in infected individuals, representing the pathogenic phase. However, few researches were found to study the mechanism of dimorphism. Thus, a diverse insight into the dimorphic switch mechanism, is urgently needed and we are the first one to research the mechanism of dimorphism.

**Methods:** Firstly, we investigated the microarray of T. marneffei in the Gene Expression Omnibus database (GEO) for differentially expressed genes (DEGs). Then Database for Annotation, Visualization and Integrated Discovery (DAVID) v6.8 was employed to analyze the underlying enrichment and pathway in biological process of DEGs. Meanwhile, protein-protein interaction (PPI) network was constructed using STRING database. On the strength of the theory that similar amino acid sequences share similar structures, which play a decisive role on the function of protein, three dimensional structures of hub-genes were predicted to further investigate the likely function of hub-genes.

**Results:** GSE51109 was elected as the eligible series for the purpose of our research, including GSM1238923 (GSM23), GSM1238924 (GSM24), and GSM1238925 (GSM25). PMAA\textsubscript{012920}, PMAA\textsubscript{028730}, PMAA\textsubscript{068140}, PMAA\textsubscript{092900}, PMAA\textsubscript{032350} were the most remarkable genes in all of the three PPI networks, thus, were viewed as hub-genes. With regard to the three-dimensional construction, except that there was no significant prediction structure of PMAA\textsubscript{092900} with the criterion seq identify > 30%, GMQE: 0-1, QMEAN4: -4-0, the parallel templates for four structures were Crystal structure of Saccharomyces cerevisiae mitochondrial NADP(+)-dependent isocitrate dehydrogenase in complex with isocitrate, Organellar two-pore channels (TPCs), Yeast Isocitrate Dehydrogenase (Apo Form) and Crystal Structure Of ATP-Dependent Phosphoenolpyruvate Carboxykinase From Thermus thermophilus HB8 in order.

**Conclusion:** The dimorphic transition of T. marneffei was viewed as a pathogenic factor and DEGs were observed. In-depth study of the function and pathway of DEGs revealed that PMAA\textsubscript{012920}, PMAA\textsubscript{028730}, PMAA\textsubscript{068140}, PMAA\textsubscript{092900}, PMAA\textsubscript{032350} were most likely acting as the hub-genes and were likely taking effect through regulating energy metabolism.

**Abbreviations:** BP = Biological Process, CC = Cellular Component, DEG = differentially expressed gene, GEO = gene expression omnibus, GO = gene ontology, GSM = GSE51109, KEGG = Kyoto Encyclopedia of Genes and Genomes, MF= molecular function, PPI = protein-protein interaction, T.marneffei = Talaromyces marneffei.

**Keywords:** bioinformatic analysis, dimorphic transition, pathogenic mechanism, talaromyces marneffei
1. Introduction

T. marneffei, previously known as Penicillium marneffei, was a crucial opportunistic thermal dimorphic fungus most prevalent in Southeast Asia\(^1\) that cause systematic mycosis, such as lung infection, bone invasion, and skin lesion. Still, there are some cases reported as endemic cases. Bamboo rats were accepted as host species of T. marneffei, and the soil of their burrows was a source of T. marneffei as well. Once infected, it can be an intractable case and is life threatening and require long-term treatment, which is costly. Over the years, an increasing number of cases occurred in immunocompetent patients have been reported, thus increased social and family burden. Phase switch, known as a factor linked with virulence, that is to say, it can grow into a multicellular hyphal form at 25°C while a unicellular fission yeast form at 37°C. In addition, in response to specific temperature cues, the hyphal form can undergo asexual development (condiation) and produce a differentiated multicellular structure called conidiophore which produces dormant spores (conidia) and inhaled conidia initiate infection and only the yeast form is found in infected individuals. Should there be a specific agent that can inhibit the binary conversion of taloromyces marneffei, less people will suffer from infection. Therefore, in this research, we pay attention to the genes of taloromyces marneffei, less people will suffer from infection. Since the 1980s, when the first case of T. marneffei infection was reported, virulence has been a fierce debate and some researches have been executed to investigate the toxicity of T. marneffei.\(^2\) The phase switch, viewed as an indispensable researches have been executed to investigate the toxicity of T. marneffei, previously known as Penicillium marneffei, was a crucial opportunistic thermal dimorphic fungus most prevalent in Southeast Asia\(^1\) that cause systematic mycosis, such as lung infection, bone invasion, and skin lesion. Still, there are some cases reported as endemic cases. Bamboo rats were accepted as host species of T. marneffei, and the soil of their burrows was a source of T. marneffei as well. Once infected, it can be an intractable case and is life threatening and require long-term treatment, which is costly. Over the years, an increasing number of cases occurred in immunocompetent patients have been reported, thus increased social and family burden. Phase switch, known as a factor linked with virulence, that is to say, it can grow into a multicellular hyphal form at 25°C while a unicellular fission yeast form at 37°C. In addition, in response to specific temperature cues, the hyphal form can undergo asexual development (condiation) and produce a differentiated multicellular structure called conidiophore which produces dormant spores (conidia) and inhaled conidia initiate infection and only the yeast form is found in infected individuals. Should there be a specific agent that can inhibit the binary conversion of taloromyces marneffei, less people will suffer from infection. Therefore, in this research, we pay attention to the genes of taloromyces marneffei closely related to dimorphism transmission, which play a key role in pathogenicity, anticipating to reveal the key factors of virulence and further explore the characteristic of them, which may contribute to find the precise agent to inhibit the growth of this fungus. Since the 1980s, when the first case of T. marneffei infection was reported, virulence has been a fierce debate and some researches have been executed to investigate the toxicity of T. marneffei.\(^2\) The phase switch, viewed as an indispensable element for virulence, the molecular mechanism of which was still unclear. Therefore, the investigations of the molecular changes are pressed for. There have been an accumulating amount of researches regarding to pathogenic mechanism of other disease, such as breast cancer and myeloid leukemia through bioinformatic tools, taking single-cell genomics, RNA sequencing, transcriptomics,\(^3,4\) proteomics\(^5\) and time resolved quantitative phospho-tyrosine analysis\(^6\) as samples, and found certain molecule to be crucial in disease progression and prognosis. Thus, bioinformatic tools were implied to research the differentially expressed genes during dimorphic switch of taloromyces marneffei and expected that some key genes would be found to be crucial in pathogenic mechanism of T. marneffei infection.

2. Materials and methods

2.1. Differentially expressed genes acquisition

The gene expression profile GSE51109, deposited by Pasricha et al.\(^7\) was downloaded from GEO database (https://www.ncbi.nlm.nih.gov/geo/). Based on the platform of the GPL17756 [Pm 5K Random Array II] Affymetrix Human Genome U133 Plus 2.0 Array, the gene expression profile contained three samples, including the genes expression with respect to intercomparing among hyphal form, yeast form and conidia. PCR products from previously cloned Penicillium marneffei genes with known expression profiles were included as controls on the microarray. Data were normalized using a global loess curve for each array\(^8\) and given normalized log2 ratio ch2/ch1 representing pair-wise combinations. Genes with log FC absolute value no less than 1 were selected as DEGs.

2.2. GO enrichment analysis and pathway analysis

Since the DEGs were screened out, the biological function and enrichment pathways of DEGs were further investigated by DAVID v6.8 (https://david.ncifcrf.gov/home.jsp), which greatly contributes to understanding biological meaning behind large list of genes by providing a comprehensive set of tools for investigators. As a consequence, several functions can be conducted with regard to the submitted gene lists, for example, identified enriched biological themes, particularly Gene Ontology (GO) terms, discover enriched functional-related gene groups, visualize genes on BioCarta & Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway maps, search for other functionally related genes not in the list, list interacting proteins and so on. Therefore, to better understand the potential function of DEGs, DAVID v6.8 was employed to analyze the underlying enrichment and pathway in biological process of DEGs.

2.3. Protein-protein interaction (PPI) network construction

As is known to us all, genes taking effect were influenced by series factors such as other genes and other micro molecule. Aiming to explore the specific relationship between DEGs, we constructed a network displaying the interaction between DEGs by STRING (https://string-db.org/cgi/aboout.pl?footer_active_subpage=content), a database which contains known and predicted protein-protein interaction, including direct (physical) and indirect (functional) associations. The interaction sources are based on computational prediction, knowledge transfer between organisms, and interactions aggregated from other databases. Genomic Context Predictions, High-throughput Lab Experiments, (Conserved) Co-Expression, Automated Text mining, Previous Knowledge in Databases make up the five main sources that give support to the interaction. To gain a deeper insight into the molecular mechanism of pathogenicity, the DEGs were uploaded to STRING then export the image of PPI and the top five most connected nodes were selected as hub-genes.

2.4. Protein three-dimensional structure construction

Genes give play to regulate the biological function or produce certain protein after transcription and translation. Thus, we were committed to predict the protein of each hub-genes through constructing the protein three-dimensional structure of each hub-genes. Accessible via the ExPASy web server, or from the program DeepView (Swiss Pdb-Viewer), SWISS-MODEL is a fully automated protein structure homology-modelling server, with the purpose of making protein modelling accessible to all life science researchers worldwide.\(^9\) Regarding to that the hub-genes were elected, further investigation into the specific functions of these genes is necessary. On the strength of the theory that similar amino acid sequences share similar structures, which play a decisive role on the function of protein, each hub-gene’s nucleotide sequence downloaded from National Center for Biotechnology Information Search database(NCBI) were submitted to NovoPro (http://www.novopro.cn/tools/translate.html) database for translation, then SWISS-MODEL was exploited for the protein three-dimensional structure construction. Afterwards, the elected templates were uploaded to RCSB-PDB.
### Table 1

The result of GO and pathway analysis of GSM23.

| Category   | Term                                      | Count | %      | P value   | Genes                                                                 |
|------------|-------------------------------------------|-------|--------|-----------|-----------------------------------------------------------------------|
| GOTERM_BP  | generation of precursor metabolites and energy | 6     | 3.0978 | 0.007248  | PMAA_039680, PMAA_028880, PMAA_046420, PMAA_068140, PMAA_089220, PMAA_071630 |
| GOTERM_BP  | inorganic cation transmembrane transport   | 4     | 2.06186 | 0.020237  | PMAA_040300, PMAA_042510, PMAA_010280, PMAA_041270                  |
| GOTERM_BP  | inorganic ion transmembrane transport      | 4     | 2.06186 | 0.020237  | PMAA_040300, PMAA_042510, PMAA_010280, PMAA_041270                  |
| GOTERM_BP  | cation transmembrane transport             | 4     | 2.06186 | 0.021917  | PMAA_040300, PMAA_042510, PMAA_010280, PMAA_041270                  |
| GOTERM_BP  | ion transmembrane transport                | 4     | 2.06186 | 0.031433  | PMAA_040300, PMAA_042510, PMAA_010280, PMAA_041270                  |
| GOTERM_CC  | mitochondrion                              | 18    | 9.27835 | 2.57E-04  | PMAA_073320, PMAA_094650, PMAA_055980, PMAA_067490, PMAA_060200, PMAA_098380, PMAA_038320, PMAA_06140, PMAA_041520 |
| GOTERM_CC  | cytoplasm                                  | 42    | 21.6495 | 0.014158  | PMAA_037440, PMAA_013400, PMAA_092900, PMAA_094610, PMAA_047150, PMAA_02080 |
| GOTERM_CC  | cytoplasmic part                           | 38    | 19.5876 | 0.01651   | PMAA_037440, PMAA_013400, PMAA_094610, PMAA_047150, PMAA_02080 |
| GOTERM_MF  | oxidoreductase activity                    | 39    | 20.1031 | 1.24E-04  | PMAA_065440, PMAA_092900, PMAA_030170, PMAA_032300, PMAA_022690, PMAA_068140, PMAA_002930, PMAA_080980, PMAA_065580, PMAA_042280, PMAA_079680, PMAA_027680, PMAA_012920, PMAA_039112, PMAA_043960, PMAA_069330, PMAA_071760, PMAA_020160, PMAA_029690, PMAA_021820, PMAA_088690, PMAA_089220, PMAA_089910, PMAA_075640, PMAA_021280, PMAA_097070, PMAA_064620, PMAA_098380, PMAA_064600, PMAA_073270, PMAA_071630, PMAA_024760, PMAA_028330, PMAA_039680, PMAA_093480, PMAA_046420, PMAA_015890, PMAA_029730, PMAA_066300, PMAA_065540, PMAA_100150 |
| GOTERM_MF  | catalytic activity                         | 99    | 51.0309 | 0.001285  | PMAA_065440, PMAA_094610, PMAA_020930, PMAA_068140, PMAA_002930, PMAA_080980, PMAA_065580, PMAA_023440, PMAA_042500, PMAA_081090, PMAA_071690, PMAA_094890, PMAA_029890, PMAA_021820, PMAA_088690, PMAA_089220, PMAA_069820, PMAA_010100, PMAA_075640, PMAA_028880, PMAA_007070, PMAA_001380 |

(continued)
As a result, the function of the protein, into which hub-genes translated can be duplicated from the templates.

3. Results

3.1. Differentially expressed genes acquisition

According to the obtained GEO series GSE51109, there are a total of 194, 169 and 157 DEGs for GSM23, GSM24 and GSM25 respectively, among which, up-regulation genes for each is 115, 98 and 80, while down-regulation genes account for 79, 71, and 77 severally.

3.2. GO enrichment analysis and pathway analysis

As is shown in Tables 1 to 3, GO enrichment was composed of biological process (BP), cellular component (CC) and molecular function (MF), taking the $P$ value of $<0.05$ as a significant criteria, the top five items of BP, CC and MF were extracted and when the significant items less than 5, all of the actual significant items were shown. In Table 1, the result of GO enrichment and pathway analysis of GSM23, namely conidiation versus hyphal cell types, were listed. As for BP, the top five items were generation of precursor metabolites and energy, orgononitrogen compound metabolic process, small molecule metabolic process, single-organism metabolic process were the top five BP stuffs. As regard to MF, isocitrate dehydrogenase activity, oxidoreductase activity, NAD binding, magnesium ion binding, cofactor binding was the top five results, while there was no result for CC and pathway analysis. The enrichment and pathway analysis results of GSM25 were displayed in Table 3, from which, we can obtain the outcome that organic acid metabolic process, membrane, dioxygenase activity, oxidoreductase activity represented for the top five items of BP, CC, MF respectively. As the same, no pathway was enriched in for the DEGs of GSM25.

3.3. PPI network construction

The aforementioned 194, 169 and 157 DEGs for GSM23, GSM24 and GSM25 were submitted to the STRING database to uncover the potential protein-protein interactions so as to get a better understanding of the potential molecular mechanism. As is shown in Figures 1 to 3, the PPI networks were constructed in response to the DEGs of GSM23, GSM24, GSM25 with the confidence score $>0.4$. According to the networks, we observed that the connections among the following genes: PMAA_012920, PMAA_028730, PMAA_092900, PMAA_064990, PMAA_032350, PMAA_016130, PMAA_059640, PMAA_036000, PMAA_082040, PMAA_080560, PMAA_019030, PMAA_082600, PMAA_083170, PMAA_007900, PMAA_068140, PMAA_042280, PMAA_006670, PMAA_040300, PMAA_093350, PMAA_071760, PMAA_029890, PMAA_002470, PMAA_021280, PMAA_060900, PMAA_073270, PMAA_071630, PMAA_024760, PMAA_040070, PMAA_093480, PMAA_066300, PMAA_100150, PMAA_037440, PMAA_013400, PMAA_092900, PMAA_076130, PMAA_086380, PMAA_094650, PMAA_067490, PMAA_041840, PMAA_079880, PMAA_045500, PMAA_071070, PMAA_012920, PMAA_043060, PMAA_015360, PMAA_020160, PMAA_082620, PMAA_010220, PMAA_089910, PMAA_039810, PMAA_054650, PMAA_081560, PMAA_055590, PMAA_064600, PMAA_079710, PMAA_065680, PMAA_052600, PMAA_059680, PMAA_028730, PMAA_015800, PMAA_032300, PMAA_014300, PMAA_022690, PMAA_020080, PMAA_002910, PMAA_027680, PMAA_091840, PMAA_093910, PMAA_039120, PMAA_063790, PMAA_021820, PMAA_009270, PMAA_092050, PMAA_064620.
dimensional structures of the five hub-genes were constructed, which were shown in Figures 4 to 7. Nevertheless, there was no significant prediction structure of the hug-gene PMAA_092900 with the criterion seq identify > 30%, GMQE: 0–1, QMEAN4: −4–0. Then the rest four significant predicted structures were confronted with the examination of the following tools: verify 3D, Procheck, Whatcheck, Errat and Prove, luckily, the above-mentioned four structures were qualified since they met the criteria of at least three inspection tools. When consult to the PDB database for the specific function of templates, representing the function of the predicted protein, the parallel templates for four structures were Crystal structure of Saccharomyces cerevesiae mitochondrial nicotinamide adenine dinucleotide phosphate (NADP)(+)‐dependent isocitrate dehydrogenase in complex with

| Category          | Term                                      | Count | %        | P value | Genes                                                                 |
|-------------------|-------------------------------------------|-------|----------|---------|----------------------------------------------------------------------|
| GOTERM_BP         | organic acid metabolic process             | 16    | 9.46756  | 5.28E-04| PMAA_080350, PMAA_069820, PMAA_005580, PMAA_028880, PMAA_097060, PMAA_068140, PMAA_046930, PMAA_012920, PMAA_013650, PMAA_005610, PMAA_039680, PMAA_031950, PMAA_046420, PMAA_089220, PMAA_028730, PMAA_041990 |
| GOTERM_BP         | generation of precursor metabolites and energy | 7     | 4.142012 | 0.01382 | PMAA_039680, PMAA_028880, PMAA_046420, PMAA_005310, PMAA_068140, PMAA_089220, PMAA_071630 |
| GOTERM_BP         | organonitrogen compound metabolic process   | 24    | 14.20118 | 0.01518 | PMAA_080350, PMAA_024190, PMAA_028110, PMAA_102650, PMAA_073320, PMAA_047150, PMAA_032300, PMAA_028880, PMAA_005580, PMAA_002470, PMAA_097060, PMAA_086380, PMAA_005310, PMAA_026340, PMAA_055500, PMAA_073270, PMAA_046930, PMAA_013650, PMAA_005610, PMAA_039680, PMAA_031950, PMAA_054240, PMAA_092720, PMAA_028730, PMAA_041990 |
| GOTERM_BP         | small molecule metabolic process            | 21    | 12.42604 | 0.01884 | PMAA_080350, PMAA_069820, PMAA_102650, PMAA_005580, PMAA_028880, PMAA_002470, PMAA_097060, PMAA_086380, PMAA_005310, PMAA_068140, PMAA_089220, PMAA_028730, PMAA_041990 |
| GOTERM_BP         | single-organism metabolic process           | 27    | 15.97633 | 0.03027 | PMAA_005310, PMAA_102650, PMAA_053530, PMAA_032300, PMAA_097060, PMAA_086380, PMAA_068140, PMAA_046930, PMAA_013650, PMAA_005610, PMAA_039680, PMAA_031950, PMAA_046420, PMAA_092720, PMAA_028730, PMAA_041990 |
| GOTERM_MF         | isocitrate dehydrogenase activity           | 3     | 1.775148 | 0.01362 | PMAA_012920, PMAA_046420, PMAA_068140 |
| GOTERM_MF         | oxidoreductase activity                     | 30    | 17.75148 | 0.011744| PMAA_005580, PMAA_065580, PMAA_068140, PMAA_042280, PMAA_079980, PMAA_057450, PMAA_018640, PMAA_027680, PMAA_012920, PMAA_043060, PMAA_031950, PMAA_071760, PMAA_029890, PMAA_005580, PMAA_046420, PMAA_035040, PMAA_012920, PMAA_005580, PMAA_046420, PMAA_035660, PMAA_065540, PMAA_022840, PMAA_100150 |
| GOTERM_MF         | NAD binding                                | 5     | 2.95858  | 0.017914| PMAA_012920, PMAA_068140, PMAA_005580, PMAA_046420, PMAA_068140, PMAA_005580, PMAA_022840, PMAA_100150 |
| GOTERM_MF         | magnesium ion binding                       | 5     | 2.95858  | 0.020596| PMAA_005580, PMAA_012920, PMAA_005580, PMAA_046420, PMAA_068140 |
| GOTERM_MF         | cofactor binding                            | 14    | 8.284024 | 0.038482| PMAA_005580, PMAA_065540 |
Table 3
The result of GO and pathway analysis of GSM25.

| Category | Term                  | Count | %      | P value | Genes                                                                 |
|----------|-----------------------|-------|--------|---------|----------------------------------------------------------------------|
| GOTERM_BP| organic acid metabolic process | 9     | 5.769230769 | .0458589 | PMAA_069820, PMAA_046930, PMAA_005610, PMAA_005580, PMAA_013650, PMAA_031950, PMAA_009220, PMAA_028730, PMAA_040360 |
| GOTERM_CC| membrane              | 38    | 24.35897436 | .0497346 | PMAA_024520, PMAA_017430, PMAA_078940, PMAA_045900, PMAA_100870, PMAA_032300, PMAA_044710, PMAA_038320, PMAA_018640, PMAA_057450, PMAA_040300, PMAA_035260, PMAA_094330, PMAA_056840, PMAA_006690, PMAA_021820, PMAA_072760, PMAA_089220, PMAA_088670, PMAA_047000, PMAA_039650, PMAA_101530, PMAA_057360, PMAA_056120, PMAA_005470, PMAA_041270, PMAA_001940, PMAA_045600, PMAA_004490, PMAA_049550, PMAA_041520, PMAA_037620, PMAA_078310, PMAA_027500, PMAA_074850, PMAA_0569270, PMAA_040940, PMAA_019030 |
| GOTERM_MF| dioxygenase activity  | 5     | 3.205128205  | .0065975 | PMAA_024760, PMAA_075640, PMAA_021280, PMAA_031950, PMAA_0397070 |
| GOTERM_MF| oxidoreductase activity| 24    | 15.38461538 | .019369 | PMAA_024520, PMAA_027300, PMAA_075640, PMAA_032300, PMAA_021280, PMAA_022690, PMAA_086460, PMAA_0097070, PMAA_005580, PMAA_057450, PMAA_018640, PMAA_024700, PMAA_075900, PMAA_005610, PMAA_039120, PMAA_021820, PMAA_031950, PMAA_082030, PMAA_009480, PMAA_021820, PMAA_089220, PMAA_006300, PMAA_028730, PMAA_022840, PMAA_100150 |

Figure 1. PPI network of GSM23. XP002153408.1, XP002144569.1, XP002146265.1, XP002148836.1 and XP002144939.1 represent the gene of PMAA-012920, PMAA-028730, PMAA-0068140, PMAA-0092900 and PMAA-032350 respectively. The size of nodes range according to the degree of analysis.
Figure 2. GSM24. XP002153408.1, XP002144569.1, XP002146265.1, XP002148836.1 and XP002144939.1 represent the gene of PMAA-012920, PMAA-028730, PMAA-0092900 and PMAA-032350 respectively. The size of nodes range according to the degree of analysis.

Figure 3. GSM25. XP002153408.1, XP002144569.1, XP002146265.1, XP002148836.1 and XP002144939.1 represent the gene of PMAA-012920, PMAA-028730, PMAA-0092900 and PMAA-032350 respectively. The size of nodes range according to the degree of analysis.
isocitrate, Organellar two-pore channels (TPCs), Yeast Isocitrate Dehydrogenase (Apo Form) and Crystal Structure Of ATP-Dependent Phosphoenolpyruvate Carboxykinase From Thermus thermophilus HB8 in order.

3.5. Validation of DEGs and hub-genenes

GSE51109 contains data about the comparisons between three phases, from which, we extracted DEGs and hub-genenes and found that PMAA_068140, PMAA_012920, PMAA_028730, PMAA_092900, PMAA_032350 were obviously differentially expressed and connected with other genes widely. In order to further verify whether their expressions were different indeed in these fungal biological states or transformation processes, we downloaded GSE51110 from the GEO database. GSE51110 gave a deep insight into the DEGs between dimorphism transition by comparing the different expression profile between hyphal cells and after transferring to 37°C as well as yeast cells and yeast cells after transferring to 25°C. Further analysis found that only PMAA_068140 was significantly different expressed according to our previous criteria. PMAA_028730, PMAA_092900, PMAA_032350 PMAA_012920 were more or less variant during dimorphic transition.

4. Discussion

T. marneffei, mostly prevalent in Southeast Asia, cause deadly systematic infection not only in immunocompromised person, but also in immunocompetent person, resulting from immuno-
supressant abuse. Once infected, it can evolve into a refractory disease. It is widely accepted that this fungus possesses the ability of dimorphic switch. And the yeast form was found in infected individuals and thus represents the pathogenic phase. Therefore, delving the potential molecular mechanism among phase transition stands a good chance to draw forth more precise therapy by molecular target therapy. Though the number of researches focuses on genome, proteome and virulence is growing year by year, there is few researches probing the expression profile difference during dimorphism transition. In present study, the DEGs in dimorphic transition were investigated for the first time and further survey their molecular function.

GO enrichment analysis shown that generation of precursor metabolites and energy was the top one for GO term of BP, and mitochondrion for CC, while oxidoreductase activity occupied the first place of MF in GSM23. This result suggested that the DEGs of condensation versus hyphal cell types played a vital part in energy metabolism. For GSM24, organic acid metabolic process, isocitrate dehydrogenase activity took the first place for GO term BP and MF respectively, which implied that the two items mentioned above were the potential mechanism in yeast versus hyphal cell types. Also, organic acid metabolic process, membrane and dioxygenase activity were the parallel terms for GSM25. Namely, the DEGs of yeast versus condensation type played a part in organic acid metabolic process, membrane and dioxygenase activity. As for pathway analysis, none of the three platforms yield significant outcome. The leading cause of this phenomenon may be the small number of elected chips, therefore, more experiments are needed to evaluate the DEGs between dimorphic transmission. To sum up, the biological function of DEGs in all the three platforms are mainly concerned with energy metabolism.

STRING functional protein association network was constructed and five genes (PMAA_012920, PMAA_028730, PMAA_068140, PMAA_092900, PMAA_032350) were found showing the strongest connections in all the three platforms surprisingly. Therefore, we have very reason to expect these five genes to be the hub-genes and their features in the switch are indispensable. What’s more, from the PPI network, we can obtain the information that the protein structures of those hub-genes remained indefinite. Thus, we predicted the three-dimensional structures of those hub-genes for further understanding of the hub-genes’ function. Except for PMAA_092900, significant three-dimensional structures of all the four genes were predicted. When referring to PDB database and PubMed database, the parallel templates for the four hub-genes: PMAA_012920, PMAA_028730, PMAA_068140, PMAA_032330 were found showing the strongest connections in all the three templates surprisingly. Therefore, we have very reason to expect these five genes to be the hub-genes and their features in the switch are indispensable. What’s more, from the PPI network, we can obtain the information that the protein structures of those hub-genes remained indefinite. Thus, we predicted the three-dimensional structures of those hub-genes for further understanding of the hub-genes’ function. Except for PMAA_092900, significant three-dimensional structures of all the four genes were predicted. When referring to PDB database and PubMed database, the parallel templates for the four hub-genes: PMAA_012920, PMAA_028730, PMAA_068140, PMAA_032330 for constructing their three-dimensional structures were extracted: Crystal structure of Saccharomyces cerevisiae mitochondrial NADP (+)-dependent isocitrate dehydrogenase in complex with isocitrate, major histocompatibility complex (MHC)-independent T-cell receptor B12A, Isocitrate Dehydrogenase (Apo Form), Crystal Structure Of ATP-Dependent Phosphoenolpyruvate Carboxykinase From Thermus thermophilus HB8. Thus, we focus on the function of those templates which may be viewed as the prediction function of the predicted protein structures.

Saccharomyces cerevisiae mitochondrial NADP (+)-dependent isocitrate dehydrogenase (NADP-IDH) in complex with isocitrate Saccharomyces cerevisiae mitochondrial NADP (+)-dependent isocitrate dehydrogenase (NADP-IDH) in complex with isocitrate plays a crucial role in Kreb’s cycle and catalyzes oxidative decarboxylation of isocitrate (ICT) into aaketoglutarate (AKG). Furthermore, it has been reported that cellular defense against oxidative damage, detoxification of reactive oxygen species, as well as synthesis of fat and cholesterol through generation of NADPH and AKG were other aspects of eukaryotic NADP-IDHs.[10] While a single type of NADP-dependent IDH contributed to most bacteria and archaea (NADP-IDH) for catalyzing reaction during the Krebs cycle, two for eukaryotes, meaning the types of IDHs employ either nicotinamide adenine dinucleotide (NAD) or NADP as a coenzyme, as Peng Y et al reported.[11] In the view of sequence analysis, the NADP-IDHs can be mainly divided into two subfamilies.[12] Subfamily II NADP-IDHs include mainly eukaryotic NADP-IDHs and a few bacterial NADP-IDHs while most bacterial and archaeal NADP-IDHs was part of subfamily I. T marneffei, accepted as a momentous fungus, being part of eukaryon, naturally, possess subfamily II NADP-IDHs and owns two type of IDHs using either NAD or NADP. Regarding to that human belongs to eukaryon as T marneffei, research into the difference of NADP-IDH between T marneffei and human being may provide a target to treat penicilliosis doing no harm to patients.

Organellar two-pore channels (TPCs) Belonging to the voltage-gated ion channel superfammy,[13] function as a homodimer with each subunit containing two homologous Shaker-like 6-TM repeats.[14] TPCs, are associated with various physiological processes in mammalian, such as autophagy regulation,[15] mTOR-dependent nutrient sensing,[16] lipid metabolism[17] and Ebola virus infection. Since TPCs are ubiquitously expressed in animals and plants[18] and TM is subjected to fungus, there was no TPCs in T marneffei. However, based on similar structure, the protein translated by PMAA_028730 may play a similar role as TPCs in fungus, that is, this kind of protein is likely linked with autophagy regulation, mTOR-dependent nutrient sensing, lipid metabolism. Nevertheless, to verify this assumption, more experiments are needed.

The yeast NAD-specific Isocitrate Dehydrogenase controlling a rate-limiting step in the tricarboxylic acid cycle by catalyzing the oxidative decarboxylation of isocitrate to α-ketoglutarate, is a hetero-octamer composed of four IDH1 and four IDH2 subunits, in which, IDH2 functions as catalytic subunit with isocitrate/Mg- and NAD-binding site, while IDH1 functions in cooperative binding of isocitrate and in binding of the allosteric activator AMP. Being consistent with yeast in the respect of microscopic species, T marneffei shares analogous functions with yeast. Thus, we can draw the conclusion that the protein translated by PMAA_068140 may act as a momentous role as NAD-specific isocitrate dehydrogenase in tricarboxylic acid cycle.

Crystal Structure of ATP-Dependent Phosphoenolpyruvate Carboxykinase From Thermus thermophilus HB8 Phosphoenolpyruvate carboxykinase (PEPCK) isoforms are well known as rate-limiting enzymes for gluconeogenesis and glycerol neogenesis pathways. Montal et al reported that inhibition of phosphoenolpyruvate carboxykinase blocks lactate utilization and impairs tumor growth in colorectal cancer.[19] Thermus thermophilus HB8, an extreme thermophile, has an optimum growth temperature of 348 K (TPePCK), which requires a more rigid thermostability. As Sugahara, M et al reported, the intra protomer hydrogen-bond interactions do not account for the thermostability of TPePCK while the entropic effects may contribute significantly to the higher stability.[20] Since TM relies on the particular temperature of 25°C or 37°C, while HB8 can withstand the extreme temperature of 348K, this predicted structure may contribute mostly to the thermostability of T marneffei.
All in all, not only in the process of GO function analysis, but also in the three-dimensional structure prediction and further investigation of specific function of each hub-gene, energy metabolism was the common ground for all of the results. Thus, we can conclude that the hub-genes were principal participating in the regulation of energy metabolism.

Still, limitations were found owing to the number of chips and bioinformatics tools elected. We aimed at revealing the secrets in the dimorphism transition, and the datasets eligible for our research was no other than GSE51109 and GSE51110, further researches about the dimorphic transition of T marneffei are calling for.

**Author contributions**

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