The de novo transcriptome of workers head of the higher group termite
*Globitermes sulphureus* Haviland (Blattoidea: Termitidae)

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**ABSTRACT**

The subterranean termite *Globitermes sulphureus* is an important Southeast Asian pest with limited genomic resources that cause damages to agriculture crops and building structures. Therefore, the main goal of this study was to survey the *G. sulphureus* transcriptome composition. Here, we performed de novo transcriptome for *G. sulphureus* workers’ heads using Illumina HiSeq paired-end sequencing technology. A total of 88,639,408 clean reads were collected and assembled into 243,057 transcripts and 193,344 putative genes. The transcripts were annotated with the Trinotate pipeline. In total, 27,061 transcripts were successfully annotated using BLASTX against the SwissProt database and 17,816 genes were assigned to 47,598 GO terms. We classified a total of 223 transcripts into COG categories, resulting in 25 groups of functional annotations. Next, a total of 12,194 genes were matched in the KEGG pathway and 392 metabolic pathways were predicted based on the annotation. Moreover, we detected two endogenous cellulases in the sequences. The RT-qPCR analysis showed that there were significant differences in the expression levels of two genes β-glucosidase and endo-β-1,4-glucanase between worker and soldier heads of *G. sulphureus*. This is the first study to characterize the complete head transcriptome of a higher termite *G. sulphureus* using a high-throughput sequencing. Our study may provide an overview and comprehensive molecular resource for comparative studies of the transcriptomics and genomics of termites.

## 1. Introduction

Termites are a group of eusocial insects found on all continents except Antarctica (Govorushko, 2018). There are about 435 species of termites in Asia (Krishna et al., 2013) and specifically over 175 species in Peninsular Malaysia (Ab Majid and Ahmad, 2011). Termites are important in the ecological ecosystem as they consume and recycle wood, litter, leaf, and dung and release the nutrients back to the ecosystem (Freymann et al., 2008; Lenz et al., 2011). Termites tunneling help in soil fertility and improvements in soil composition (Govorushko, 2018).

Despite the status as ecological engineers, a few species of termites are regarded as pests, infesting and damaging structural woods of buildings, houses as well as agricultural plantations (Kuswanto et al., 2015). Damages caused by termites has economic impact, resulting in 25 groups of functional annotations. Next, a total of 12,194 genes were matched in the KEGG pathway and 392 metabolic pathways were predicted based on the annotation. Moreover, we detected two endogenous cellulases in the sequences. The RT-qPCR analysis showed that there were significant differences in the expression levels of two genes β-glucosidase and endo-β-1,4-glucanase between worker and soldier heads of *G. sulphureus*. This is the first study to characterize the complete head transcriptome of a higher termite *G. sulphureus* using a high-throughput sequencing. Our study may provide an overview and comprehensive molecular resource for comparative studies of the transcriptomics and genomics of termites.

The subterranean termite, *Globitermes sulphureus* (Haviland) (Blattoidea: Termitidae), is a higher wood-feeding termite that is distributed throughout Southeast Asia, including Singapore, Thailand, and Malaysia (Hussin et al., 2018). This species can be easily recognized by the bright yellow colored abdomen of its soldier (Lee et al., 2002). *G. sulphureus* is an important agricultural pest, especially in coconut and oil palm plantations and is also regarded as a secondary pest where the members of this species infest premises that have been treated with termite bait (Bakaruddin et al., 2018; Lee et al., 2007; Neoh et al., 2011). *G. sulphureus* is commonly found infesting doors, window frames, wooden walls, wooden pillars, and carpet grippers (Ab Majid and Ahmad, 2009; Ngee and Lee, 2002).

Although *G. sulphureus* has a significant economic impact, its genomic data in public database are particularly scarce and only a few sequences have been published. The available sequences are mainly corresponded to mitochondrial (Bourguignon et al., 2017), ribosome fragments (Yap et al., 2007), and endo-β-1,4-glucanase (Bujang et al., 2014). Even for other termites species, to date only three termite genome sequences are

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publicly available: two from the lower termite Zootermopsis nevadensis (Terrapon et al., 2014) and Cryptotermes secundus (Harrison et al., 2018) and one from the higher termite Macrotermes natalensis (Poulsen et al., 2014). The genome size of Z. nevadensis is 562 Mb smaller than C. secundus (1.30 Gb) and M. natalensis (1.31 Gb) (Harrison et al., 2018) while the cockroach Blattella germanica genome (2.0 Gb) is considerably larger than all three termite genomes.

The actual genome size of G. sulphureus is unknown but a study done by Koshikawa et al. (2008) has mentioned that the Termitidae genome size based on C-value ranged from 1.48 pg to 1.90 pg. The rapid development of next-generation sequencing (NGS) technologies offers the opportunity to gather population genomic data in non-model organisms, in the absence of prior knowledge, at an affordable cost (Gayral et al., 2013). RNA sequencing (RNA-Seq) has been successful and increasingly used to define the transcriptome in termites (Dedeine et al., 2015; Geng et al., 2018; Huang et al., 2012; Rosenthal et al., 2011; Yuki et al., 2008).

Genomes of higher termites are considered large, highly repetitive and, consequently, difficult to assemble (Korb et al., 2015). We therefore, decided to assemble its transcriptome. De novo transcriptomics (assembling transcriptomes without a reference genome) gives access to large numbers of genes at relatively low cost, with potential applications for SNP discovery and molecular markers (Gayral et al., 2013; Khizam and Ab Majid, 2019; Levy and Myers, 2016). De novo transcriptome has also been applied in other insect studies such as beetle (Sayadi et al., 2016), ants (Morandin et al., 2018), and cockroach (Kim et al., 2016).

We have previously recorded the metagenomic data from the worker and soldier gut of G. sulphureus (Hussin et al., 2018) and isolated their gut fauna using culture-dependent method (Hussin and Ab Majid, 2017). Lower and higher group termites were proven synergistically with their gut symbionts for lignocellulose digestion (Franco Cairo et al., 2016; Ni and Tokuda, 2013). Termites and their gut symbionts produced various cellulase enzymes in the salivary gland, midgut, and hindgut (Bruno, 2014; Ni and Tokuda, 2013). These enzymes have demonstrated great potential for bioenergy applications (Bruno, 2014; Scharf and Tartar, 2008; Singhania et al., 2013). Endo-β-1,4-glucanases and β-glucosidases are the key enzymes for cellulose digestion (Bruno, 2014; Bujang et al., 2014; Lima et al., 2014; Ni and Tokuda, 2013).

In this present study, we used RNA-Seq approach to transcriptome profiling of worker heads from G. sulphureus using the Illumina HiSeq sequencing machine. The RNA was extracted from the head in order to minimize contamination by sequences from endosymbionts that inhabit the sequencing machine. The RNA was extracted from the head in order to minimize contamination by sequences from endosymbionts that inhabit the

2. Materials and methods

2.1. Termite materials for RNA-seq

The G. sulphureus colony was collected from a mound at the main campus of Universiti Sains Malaysia, Penang, Malaysia during August 2017. The termites together with the nest were transported to the laboratory using the black bucket and were used instantly. Only healthy workers (50 individuals) were selected from this single colony. The worker heads were separated from the bodies by using dissecting scissors. Then these 50 heads were pooled together and immediately stored in RNAlater RNA Stabilization Reagent (Qiagen, Germany) for further processing.

2.2. RNA extraction, cDNA library construction, and illumina sequencing

The head sample was shipped in dry ice to Major Biosystem Corporation, Taiwan for RNA extraction, cDNA library construction, and Illumina sequencing. Total RNA was extracted using TRizol® reagent (Invitrogen, USA) according to the manufacturer’s protocol. RNA purity was checked using a NanoDrop 2000 spectrophotometer. The RNA integrity was observed on 1% agarose gel electrophoresis and RNA integrity value was analyzed using Agilent 2100 Bioanalyzer. The sample was verified to have a RIN (RNA integrity number) value of 7.4. The RNA was used for the building of cDNA libraries using the Truseq™ RNA sample prep Kit (Illumina, San Diego, USA) as recommended by the manufacturer. The cDNA was amplified by PCR using Phusion DNA polymerase (NEB) for 15 PCR cycles and cDNA Library was obtained by gel extraction selected for target fragments on 2% Low Range Ultra Agarose (Certified Low Range Ultra Agarose, Bio-Rad). The library was then quantified using TBS380 (Picogreen, Invitrogen, Carlsbad, CA, USA) followed by bridge PCR amplification using the eBot Cluster Generation System (Truseq PE Cluster Kit v3-eBot-HS, Illumina, San Diego, USA). At last, the cDNA library was sequenced on the Illumina HiSeq X10 platform which generated 2 × 150 bp of paired-end raw reads.

2.3. De novo transcriptome assembly

The raw reads (FASTQ data) were quality checked for base sequence quality, GC content, base N content, and adapter content using FastQC v0.11.5 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc). These raw reads were then trimmed using Trimmmomatic software v0.36 (Bolger et al., 2014) to remove adapters, low-quality reads, and to discard the reads with a minimum length of 36 bp. The resulted sequences were de novo assembled using Trinity software v2.5.1 with a default k-mer size of 25 (Grabherr et al., 2011; Haas et al., 2013) following default protocols. The outputs were saved in the FASTA format. The quality of the assembled transcripts has been verified using Bowtie2 v2.3.3.1 (Langmead and Salzberg, 2012).

2.4. Data deposition

All sequences acquired in this research were deposited under accession number (SRP147368) connected with BioProject PRJNA471675 in the National Center for Biotechnology Information (NCBI) Sequence Read Archive (SRA) databases.

2.5. Transcriptome function annotation

The all assembled transcripts were functionally annotated using Trinity and Blastp. Blastp was applied to the assembled sequences to search for nucleotide sequences homologous using BLASTX (NCBI-blast v2.7.1+) against several databases (SwissProt database, Kyoto Encyclopedia of Genes and Genomes (KEGG), Gene Ontology (GO), and evolutionary genealogy of genes: Non-supervised Orthologous Groups (EggNog). The transcripts were also searched for potential ribosomal RNA using Rfam (v1.2) (Lagesen et al., 2007). The transdecoder peptide sequence file for final candidate longest-ORF was searched for amino acid sequence homologies using BLASTP (NCBI-blast v2.7.1+) protein domain identification via a Pfam search using Hmmer v3.1b2 software (Finn et al., 2011), presence of signal peptide cleavage sites using SignalP v4.1 (Petersen et al., 2011), and prediction of transmembrane helices in proteins using TMHMM v2.0c (Krogh et al., 2001).

All the results above were loaded into the Trinotate SQLite database and reported in Excel format. The GO annotations were extracted from
the Trinotate output and were visualized using the Web Gene Ontology Annotation Plot (WEGO) 2.0 for plotting the biological process, molecular functions, and cellular components of GO terms (Ye et al., 2006). The Cluster of Orthologous Groups (COG) functional annotations were extracted from Trinotate output which derived from the EggNog database.

2.6. Validation and relative gene expression using real-time quantitative PCR

Total RNA was extracted from the heads of worker (30 heads) and soldier (30 heads) castes using a combination of Trizol reagent (Invitrogen, USA) and with RNeasy® MinElute® Cleanup Kit (Qiagen, Germany) following to manufacturer's protocol. The total RNA was treated with RNase-free DNase (Qiagen, Germany) to remove any trace of genomic DNA. The treated RNA samples were converted into cDNA using SuperScript™ III First-Strand Synthesis (Invitrogen, USA). The cDNA products were then used as a template in real-time qPCR. The RT-qPCR mixture was prepared using iTaq™ Universal SYBR® Green Supermix (Bio-Rad, USA). The reaction mixture (10 μL) contained 5 μL of 2X iTaq™ Universal SYBR® Green Supermix, 0.5 μL of each forward and reverse primers, 1 μL of cDNA template, and 3 μL of DEPC-treated water. The RT-qPCR was performed with the Bio-Rad® CFX96™ machine with the following conditions: polymerase activation at 95 °C for 30 s, followed by 40 cycles of DNA denaturation at 95 °C for 5 s and annealing/extension + plate read at 60 °C for 30 s, at last, melt-curve analysis at 65 °C - 95 °C with 0.5 °C increment/step. Two interesting genes were selected for validation: β-glucosidase and endo-β-1,4-glucanase. Primer sequences were listed in Table 1. Data generated from RT-qPCR were analyzed using the 2^{-ΔΔCT} method (Livak and Schmittgen, 2001). All qPCR has been repeated in the three biological and three technical replications. Two genes expression was normalized against β-actin expression. Differences in the expression level of the two genes between workers and soldiers were tested using the t-test.

3. Results

3.1. Illumina sequencing and de novo assembly

In total, 93, 347, 116 raw sequencing reads were generated using the Illumina HiSeq X10 sequencing platform. Table 2 summarizes the statistics of the sequencing data and Trinity de novo assembly. After data filtering with Trimmomatic software, a total of 88, 639, 408 (95%) high-quality reads were obtained and used for assembly. De novo assembly using Trinity results in 243, 057 of transcripts and 193, 344 of genes with a GC percentage of 40.43%. The average length of transcripts was 813 bases and an N50 length was 1581. The quality of the assembled transcripts was checked using Bowtie2 and the result was 97% of the were correctly assembled.

3.2. Transcriptome annotation

Among 243, 057 Trinity transcripts, 27, 061 (11.13%) had top BLASTX hits in the SwissProt database. Meanwhile, only 51, 655 (21.25%) of all transcript sequences were effectively translated into peptide sequences using TransDecoder. From these numbers, 34, 893 (67.55%) of peptide sequences had top BLASTP hits in the SwissProt database and matched to 32, 221 (62.38%) protein entries in the Pfam database.

3.3. Functional classification by GO

In total, 17, 816 genes were assigned to 47, 598 GO terms. The GO terms were assigned to cellular compartment (16, 139, 33.91%), molecular function (15, 954, 33.52%), and biological process (15, 505, 32.57%). The terms were derived from 66 different functional groups (GO sub-categories level 2) (Figure 1). Under cellular compartment category, cell (14, 763 genes, 82.9%), cell part (14, 724 genes, 82.6%), and organelle (12, 020 genes, 67.5%) represented the majorities categories (Figure 1). For molecular function categories, binding (12, 855 genes, 63.4%) were the dominant subcategories.

3.4. Functional classification by COG

In total, 14, 223 transcripts were assigned to multiple COG classification, resulting in 16, 211 of functional annotations. The results showed that these sequenced were annotated to 25 COGs functional categories (Figure 2). Among those categories, “General function prediction only” (2, 999, 18.5%) represented the largest group, followed by “Signal transduction mechanisms” (1, 560, 9.62%), “Posttranslational modification, protein turnover, chaperones” (1, 384, 8.54%), and “Translation, ribosomal structure and biogenesis” (1, 367, 8.43%). A few other categories, such as “Cell motility”, “Mobiome: prophages, transposons”, “Nuclear structure” were the smallest groups and “Extracellular structures” categories were absent.

3.5. Functional classification by KEGG

As a result, 12, 194 genes were matched in the KEGG pathway and were assigned to 392 KEGG pathways. The distribution of KEGG pathways was shown at the second hierarchical stage (Figure 3). The top five

Table 1. Primers used in the relative gene expression experiment.

| Primers           | Sequence 5'-3'            |
|-------------------|---------------------------|
| β-glucosidase F   | CCAACAGTTAATCCTCACCTAT    |
| β-glucosidase R   | GACACCAACAGGTTCTTTACT     |
| Endo-β-1,4-glucanase F | CAGGACTTGACAGGGGATA       |
| Endo-β-1,4-glucanase R | TAGTCCTGAGGGCTATTAC       |
| β-actin F         | CTCTTCAGCTCAGCTCCTCT      |
| β-actin R         | CTCTTGATCATCCGTCAGCAA      |
largest pathway groups were signal transduction (1439, 11.80%), global and overview maps (1346, 11.04%), infectious disease: viral (694, 5.69%), endocrine system (680, 5.58%), and cancers: specific types (526, 4.31%).

3.6. Gene expression analysis

To validate the RNA-seq outcomes of the worker head samples, two cellulase genes were selected, β-glucosidase and endo-β-1,4-glucanase. The relative gene expression experiment was performed using the RNA sample derived from the worker and soldier heads. Figure 4 shows the relative expression levels of both cellulases in both castes. The β-glucosidase was considerably greater in the head of the soldiers compared to the workers ($p < 0.05$). While the endo-β-1,4-glucanase expression level in worker heads was significantly higher than soldiers ($p < 0.05$).

4. Discussions

In the present study, we characterized the transcriptome data from the worker heads of the higher termite, *G. sulphureus* using the Illumina
Figure 3. KEGG orthology (KO) classification. The second hierarchical stage of KEGG pathways were assigned into six categories: metabolism, genetic information processing, environmental information processing, cellular processes, organismal systems, and human diseases.
HiSeq sequencing platform. Several transcriptome studies in termite species have been carried out by two of which have sequenced the worker heads from *Coptotermes gestroi* (Leonardo et al., 2011) and *Odontotermes formosanus* (Huang et al., 2012). While the rest of the studies have focused on the host termite, gut microbial symbionts, or combined the host and symbionts (Scharf, 2015). The transcriptome of *G. sulphureus* was not recorded before, so this is the first effort to examine the head transcriptome of this species.

The GO describes the functions of gene products in terms of their association with cellular components, molecular functions, and biological processes in a species-independent manner (Xie et al., 2012). In the cellular component category, most of our gene products were located in the cell and cellular part component. These genes were mainly molecular functions as binding and catalytic activity that were involved in the biological process category which was cellular processes and metabolic processes. These results represent most components are related to each other and important for maintaining the basic functions of genes in the cell and tissues of an organism. These components were also abundantly found in not only termites *O. formosanus* (Huang et al., 2012), *C. gestroi* (only in the biological process) (Leonardo et al., 2011), dry wood termite *Cryptotermes domesticus* (Wu et al., 2015), but also in other insects such as seed beetle *Callosobruchus maculatus* (Sayadi et al., 2016), oriental armyworm *Mythimna separata* (Bian et al., 2017), pear *Pyrus calleryana* Decne (Xu et al., 2015), and purple sweet potato *Impomoea batatas* L. (Xie et al., 2012).

The Cluster of Orthologous Groups (COG) is a database where the proteins are classified into 26 functional categories (A to Z) based on the orthology concept (Tatusov, 2000; Tatusov et al., 2003). From the results, the generated transcriptome profiles for the heads of *G. sulphureus* were not similar to *O. formosanus* except that both termites have the “general function prediction only” as the largest group in the COG categories (Huang et al., 2012). The “general function prediction only” is a member of a poorly characterized group where most of the proteins have similar biochemical activity but different in functions (R. L. Tatusov, 2000). The same goes for the KEGG pathways where both termites have a different pattern of pathways classification. The different diet of these termites (*O. formosanus*: fungus-feeding termite and *G. sulphureus*: wood-feeding termite) and distantly related in a phylogenetic relationship (Bujang et al., 2014) may contribute to the dissimilarity in transcriptome profiles.

Two cellulase genes homologous to β-glucosidase and endo-β-1,4-glucanase were selected to validate the RNA-seq. Both genes also were selected to compare their expression levels between workers and soldiers. The expression of β-glucosidase was considerably greater in soldiers, while endo-β-1,4-glucanase was significantly higher in workers (Figure 4). These results were contradicted with the previous study in which *C. gestroi* workers' caste had a high-level expression of β-glucosidase compared to the caste of soldiers (Leonardo et al., 2011). A study of *Nasutitermes corniger* exhibited the greater level of exoglucanase and β-glucosidase activities in soldiers (Lima et al., 2014). Their relative expression analysis and enzyme activities were using the whole body or whole gut of the workers and soldiers, respectively. It is important to note that the relative expression level of genes does not correspond with its activity in the gut (Fujita et al., 2008). Further investigation concerning the digestive system of *G. sulphureus* should focus on enzyme activities for each host and gut compartment.

Since we prepared cDNA libraries from the worker and soldier heads, the cellulases we identified were likely expressed in the salivary gland. This finding was consistent with previous studies that β-glucosidase and endo-β-1,4-glucanase were found expressed in *C. gestroi* (Leonardo et al., 2011), *Cryptotermes domesticus*, *Coptotermes formosanus*, *Ahmaditermes sichuanensis* (Li et al., 2012), and *Reticulitermes flavipes* (Tartar et al., 2009). From the nucleotide sequence homology using BLASTX, our β-glucosidas and endo-β-1,4-glucanase have been categorized into GH1 and GH9 (glycoside hydrolases) families. GH1 and GH9 are groups of enzyme families generated by the termite itself rather than the gut symbionts. This was consistent with the past studies of lower termites (Franco Cairo et al., 2016; Tartar et al., 2009) and higher termites (Bujang et al., 2014).

Cellulose digestion into glucose involves three different enzymes: endo-β-1,4-glucanase (EC.3.2.1.4), exoglucanase or cellulohydrolases (EC.3.2.1.91), and β-glucosidases (EC.3.2.1.21) (Li et al., 2012). Endo-glucanases cut at random glycoside bonds within amorphous cellulose chains, while cellulohydrolases cleave at the non-reducing ends of crystalline cellulose producing cellobiose, glucose, and cellotriose. The β-glucosidas (cellobiases) then hydrolyze the cellobiose to glucose (Lima et al., 2014; Terra and Ferreira, 1994). Workers and soldiers have different feeding habits, and the results obtained from this study may reflect the biological differences between castes. The worker caste forages and feeds other colony members. The soldier caste defends the colony from enemies and is fed by worker caste via trophallaxis. Thus, they receive the partially digested food from workers such as regurgitated food, saliva, or liquid excreta rich in symbionts (Nalepa, 2015). Lima et al. (2014) mentioned that the food originating from lignocellulose products requires the enzyme endo-β-1,4-glucanase for partial digestion. As a result, the expression of this enzyme is high in workers. This partially digested food is then supplied to the soldiers engaged in the enzymes exoglucanase and β-glucosidase for the final digestion of the cellulose component, resulting in a high-level expression of β-glucosidase in soldiers.

5. Conclusions

Our data transcriptome provided the functional annotations of genomic and molecular elements in the cells and tissues of *G. sulphureus*. These annotations can be used as a reference for comparative studies and genomic analysis of termites in the future. From our qPCR outcomes,
further in-depth molecular studies and enzyme operations of two cellulase genes should be performed to elucidate the metabolic pathways engaged in lignocellulose enzymes and their possible use in biotechnological procedures.

Declarations

Author contribution statement

Nurul Akmar Hussin: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Wrote the paper.

Mohd Nazalan Mohd Najimuddin: Conceived and designed the experiments; Contributed reagents, materials, analysis tools or data.

Abdul Hafiz Ab Majid: Conceived and designed the experiments; Contributed reagents, materials, analysis tools or data; Wrote the paper.

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Competing interest statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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