Database Update

*Tetrahymena* Functional Genomics Database (TetraFGD): an integrated resource for *Tetrahymena* functional genomics

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The ciliated protozoan *Tetrahymena thermophila* is a useful unicellular model organism for studies of eukaryotic cellular and molecular biology. Researches on *T. thermophila* have contributed to a series of remarkable basic biological principles. After the macronuclear genome was sequenced, substantial progress has been made in functional genomics research on *T. thermophila*, including genome-wide microarray analysis of the *T. thermophila* life cycle, a *T. thermophila* gene network analysis based on the microarray data and transcriptome analysis by deep RNA sequencing. To meet the growing demands for the *Tetrahymena* research community, we integrated these data to provide a public access database: *Tetrahymena* functional genomics database (TetraFGD). TetraFGD contains three major resources, including the RNA-Seq transcriptome, microarray and gene networks. The RNA-Seq data define gene structures and transcriptome, with special emphasis on exon–intron boundaries; the microarray data describe gene expression of 20 time points during three major stages of the *T. thermophila* life cycle; the gene network data identify potential gene–gene interactions of 15,049 genes. The TetraFGD provides user-friendly search functions that assist researchers in accessing gene models, transcripts, gene expression data and gene–gene relationships. In conclusion, the TetraFGD is an important functional genomic resource for researchers who focus on the *Tetrahymena* or other ciliates.

Database URL: http://tfgd.ihb.ac.cn/

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**Introduction**

*Tetrahymena thermophila* is a free-living ciliated protozoan that normally has two types of functionally distinct nuclei (1), the silent germ line micronucleus (MIC) and the actively transcribed somatic macronucleus (MAC) in each cell. Its typical eukaryotic biology and many molecular genetic tools have enabled *Tetrahymena* researchers to contribute to landmark discoveries of fundamental eukaryotic cellular mechanisms, such as the first cytoskeletal motor (2), catalytic RNA (3), telomere structure (4) and telomerase (5) and the role of small RNAs in programmed somatic genome rearrangement (6). Although its genome has been sequenced and its genetics and molecular biology have been extensively studied, research on *T. thermophila* has been limited by the lack of some basic genomic resources, in particular, functional genomics data.
In 2006, the MAC genome sequence project of *T. thermophila* was completed (7), which provides the first ciliate genome sequence. After this, the *Tetrahymena* genome database (TGD and TGD Wiki, http://ciliate.org), containing the genome sequence and the predicted gene models, was established (8–9). The first insights into the *Tetrahymena* transcriptome came from expressed sequence tag (EST) sequences generated by the initial genome sequence project and the protist EST program, and they could be accessed through the NCBI EST database and the PEP database (TBESTDB, http://www.bch.umontreal.ca/pepdb/pepdb.html), respectively. Miao et al. (10) initiated *Tetrahymena* functional genomics by establishing the first *T. thermophila* microarray platform performing a genome-wide investigation of gene expression during the three major physiological and developmental stages of the *T. thermophila* life cycle. These microarray data could be accessed via the *Tetrahymena* gene expression database (TGED, http://tged.ihb.ac.cn/) (11). Recently, the microarray data were extended to infer a *T. thermophila* gene network (12), providing the first insight of gene–gene relationships in *T. thermophila*. Finally, an analysis using deep RNA sequencing (RNA-Seq) provided more comprehensive and detailed analysis of the transcriptome and greatly improved the gene models (13). Given these recent genomic studies, an integrated database is needed to provide easy access to functional genomics data for members of the scientific research community interested in *T. thermophila*.

In this study, we describe the *Tetrahymena* functional genomics database (TetraFGD), which provides user-friendly search functions for accessing gene models, transcripts, gene expression data and gene–gene relationships in *Tetrahymena*.

**Content**

The TetraFGD is an online resource containing three major functional genomic data sets of *T. thermophila*: (i) RNA-Seq data; (ii) microarray data; and (iii) gene network.

**RNA-Seq data**

These data were obtained from six RNA samples in three major physiological and developmental stages of *T. thermophila,* including one in growth (3.5 ± 10^5 cells/ml), three in starvation (mating type V and VI in 3 h, mating type VI in 15 h) and two in conjugation (2 and 8 h after mixing of two mating types) by using Illumina deep RNA sequencing (13). To ensure that the RNA-Seq data could be compared with the microarray data, time points were selected a subset of those in the previous microarray expression studies that covered 20 states of the three stages of the *T. thermophila* life cycle. More than 96% of the predicted genes have detectable reads after mapping the RNA-Seq data to the genome. More than 30 000 transcripts were assembled, including >1000 new transcripts, which were not found by gene scanning. These transcripts were used to improve the previous gene models. Over 7000 predicted gene models showed errors when the RNA-Seq data and gene annotation (13) were compared, greatly improving identification of coding sequences, untranslated regions and exon–intron boundaries. Although the RNA-Seq technology is powerful, some assembled transcripts do not contain complete open reading frames because of limited coverage and assembly. Compared with the RNA-Seq, gene scanning may be relatively less accurate when it was used to predict the transcription information, whereas it supplies the complete open reading frames. Thus, we shared the RNA-Seq data with the *Tetrahymena* team at the Broad Institute, who are using them to update gene prediction and genome annotation.

**Microarray data**

The TetraFGD now contains the microarray gene expression data containing 20 time points during the three major physiological and developmental stages of the *T. thermophila* life cycle, including 3 points in growth, 7 points in starvation and 10 points in conjugation. Because the microarray expression values might be wrong if a gene was mis-predicted, and the RNA-Seq assembled transcripts provide more correct transcription information. Therefore, the TetraFGD now integrates two types of microarray expression values: (i) normalization based on the predicted genes (gene model may be incorrectly predicted); and (ii) normalization based on the RNA-Seq assembled transcripts (some of them are transcription fragments). These two types of microarray expression values were both normalized using the microarrays reported previously (10).

There are two kinds of gene expression values normalized by the predicted genes. Take the expression values of the gene THERM_00257230 (http://tfgd.ihb.ac.cn/search/detail/gene/THERM_00257230) for example, two expression profile were showed when you search the database and are represented by the blue and red line. The blue line represents the expression values from the *Tetrahymena* gene expression database (TGED) (11), and the red line represents the expression values that were normalized by Prof. Ronald Pearlman laboratory using a different method (most of the raw data are the same as those used in the blue line, and 10 new microarrays in the *Tetrahymena* conjugation stage were added).

For the RNA-Seq assembled transcripts (each transcript was regarded as a gene model, although it may be only a transcription fragment), all previous designed microarray probes (10) were re-mapped to the RNA-Seq assembled transcripts, and the microarray expression values for these transcripts were re-normalized as in the study by Miao et al. (10) and provided in the TetraFGD.
Gene network
Network analysis can be used to identify genes in the same biological processes or pathways, to infer interactions of bio-molecules, such as their physical association, metabolite flow, regulatory relationships and co-expression relationships and so forth. An important resource in the TetraFGD is the *Tetrahymena* gene network (TGN). The TGN was constructed using the context likelihood of relatedness algorithm (CLR, mutual information-based method, which is an extension of the relevance networks) (14) based on 67 Roche NimbleGen single-channel microarray expression data (12), which means that the connected genes in the TGN have similar expression profiles. After gene filtering strategies (12), 15,049 genes were used to infer the gene network. By determining an appropriate threshold with a CLR Z-score threshold 3.49, 1,958,477 gene–gene interactions were included in the TGN (12). The larger the Z-score value between the two genes in the TGN, the more reliable interaction (more similar expression profile) between them. Several experimentally verified cases showed that the TGN-predicted gene connections were likely to have related functions, such as the proteasome complex (12), the adenosine triphosphate synthase complex (15) and genes involved in DNA rearrangement during *Tetrahymena* MAC development (6). Thus, the TGN presents an important resource to study *Tetrahymena* genes at the pathway level.

Construction
The schema of the TetraFGD is showed in Figure 1. Gene models, transcript sequences, microarray expression data and gene network data were stored in the MySQL database, and transcript sequences were formatted as a Basic Local Alignment Search Tool (BLAST) database. The web interface for searching these data was written by using Hypertext Preprocessor (PHP). Apache2 (http://httpd.apache.org/) in a CENTOS operating system was used as the web server. To provide a convenient way to check the gene models and the transcripts, Gbrowse2 (http://gmod.org/wiki/GBrowse) was set-up for graphically viewing the RNA-Seq data. A BLAST web server was also applied for the sequence-based searching against the formatted transcript database. Search functions make it easy to access to the three major resources in the TetraFGD.

Use and discussion
The TetraFGD can be accessed via the World Wide Web at http://tfgd.ihb.ac.cn/. An integrated searching box was designed to provide quick access to the data on the top of each page of the TetraFGD website. You can use ‘Gene ID’, ‘Keyword’ or ‘Transcript ID’ to search the database (Figure 2A). If you use the ‘Gene ID’ to search (Figure 2A), you should provide a ‘TTHERM_XXXXXXXX’ style ID, which was originally generated by gene prediction, and this style ID is also used in the *Tetrahymena* genome database (TGD, http://www.ciliate.org/). If you use a ‘Keyword’ to search (Figure 2A), you can type any word(s) in the search box, and the database will return the record(s) with a gene name (based on the gene annotation, but not the gene name from individual published studies) containing your keyword. If you use the ‘Transcript ID’ to search (Figure 2A), you should provide a transcript fragment ID from the RNA-Seq database. This ID can be found by either of two ways: (i) using the ‘TTHERM_XXXXXXXX’ to find its related transcript fragment ID(s) through searching the database or the look-up table (http://tfgd.ihb.ac.cn/index/version) and (ii) using your sequence to BLAST against the RNA-Seq assembled transcript database (http://tfgd.ihb.ac.cn/tool/blast).

In addition to the integrated search box, we have also designed the individual searching function, such as RNA-Seq (Figure 2B), microarray (Figure 2C) and gene network (Figure 2D).

Searching RNA-Seq data
The TetraFGD displays the RNA-Seq data graphically and performs its search function through Gbrowse (http://gmod.org/wiki/Main_Page). Typically, Gene ID can be used to search the data (Figure 2B), and it also accepts a keyword (gene annotation), a transcript ID or a scaffold region. On the Gbrowse search result page, four tracks are shown, including a predicted gene model track, a RNA-Seq assembled transcript track (linked to the transcript sequence and its microarray expression information), a RNA-Seq coverage plot track and a microarray probe track (Figure 3A). Through these tracks, you can check whether there are any gene prediction mistakes, and retrieve the transcript sequence and the gene exon–intron structure information. These data are useful for studying downstream gene function. Through the Gbrowse, you can choose any specific interesting region and export the FASTA format sequence of any selected region by clicking the ‘Download Decorated FASTA File’ in the pull-down box. In addition, Gbrowse also allows the user to export the high-resolution image.

Searching microarray data
TetraFGD currently contains the microarray data of 20 time points during the three major physiological and developmental stages of the *T. thermophila* life cycle. Gene ID can be used to search the data either in the top search box or on the individual microarray search page (Figure 2C). You can search the microarray expression values either based on the predicted gene models or the RNA-Seq assembled transcripts. For the predicted gene models, the microarray result page follows the style of the TGED, and it also
Figure 1. The schema of the TetraFGD.

Figure 2. Screenshots of searching function interfaces of the TetraFGD website. (A) Integrated searching box. The red arrow indicates the pull-down button to the searching menu for choosing the 'Gene ID', 'Keyword' or 'Transcript ID'. (B–D) Individual searching boxes for RNA-Seq, microarray and gene network.
includes the Gene ID (hyperlink to the TGD), description, sequence (cDNA and protein) and shows the expression profile [see detail in (11)]. For the RNA-Seq assembled transcripts, re-normalized microarray expression data were added to the TetraFGD to help researchers get more accurate expression information, and the result gives the summary information, sequence, as well as the expression profile (Figure 3B). In addition, a new function was designed to retrieve the detailed gene expression values for each state by clicking the button ‘Value Table=’ (Figure 3B).

**Searching Tetrahymena gene network data**

TGN was constructed with 15,049 genes (12) using the CLR algorithm (14), and it supplies a source to retrieve possible functionally related genes. On the gene network page, one can enter a gene ID for ‘Single-Search’ and multiple gene IDs for ‘Multi-Search’ (Figure 2D). These two types of search functions were designed for different needs. The ‘Single-Search’ was designed to find all the potential interacted candidates of the query gene, and the result will return a list of candidate genes with expression patterns similar to your query gene in the TGN (Figure 3C). The ‘Multi-Search’ was designed to find the potential interactions among a set of genes, and the result will return gene–gene interactions among your supplied genes in the TGN (Figure 3D). All the search results can be downloaded as a tab-delimited text file that can be visualized by using the Cytoscape software (http://www.cytoscape.org/) (16).

**Other services in the TetraFGD**

Besides the previously described functions, the TetraFGD now set-up a BLAST web server (17), allowing the RNA-Seq assembled transcripts to be found using the nucleotide sequence by BlastN or the protein sequence by TBLASTN (http://tfgd.ihb.ac.cn/tool/blast). Moreover, detailed ‘Help’ information is provided to assist in using the database easily. On the TetraFGD ‘Search Help’ page (http://tfgd.ihb.ac.cn/index/schhelp), we have supplied detailed introductions and explanations of search functions to access to all the resources. The page ‘Sample Preparation’ (http://tfgd.ihb.ac.cn/index/smphelp) describes the standardization of culture conditions of growth, starvation and conjugation, used in preparing RNA for microarrays and RNA-Seq.

It is worth noting that *T. thermophila* gene annotations have been updated several times. The microarray platform was designed according to the 2006 version genome annotation, whereas the RNA-Seq data were analysed using the 2008 version genome annotation. There are some differences between these two versions of genome annotation. Therefore, we provide a look-up table (http://tfgd.ihb.ac.cn/index/version) for the conversions among 2006 version genome annotation IDs, 2008 version genome annotation IDs, transcript IDs and gene descriptions to conveniently use this database. The user will receive a warning when the search uses an improper gene ID.

**Further development of the TetraFGD**

Future development of the TetraFGD will include uploading and integrating additional *Tetrahymena* functional
genomics data sets, such as the microarray gene expression, transcriptome, re-sequencing data of *T. thermophila* under stress from exposure to pollutants or displaying the effects of specific gene mutations and also the phosphorylation proteomics.

**Conclusions**

The TetraFGD website makes substantial improvement to the original TGED website through the addition of databases of RNA-Seq and gene network as well as BLAST searching. To facilitate access to these resources, a user-friendly web interface was developed. The TetraFGD website (also the earlier TGED website) has already attracted considerable interest from the worldwide scientists, and web traffic records indicated that they receive, on average, >150 unique visits from >40 countries per day during the past 2 years. In conclusion, the TetraFGD is an important integrated functional genomics resource, which is freely available to interested researchers.

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**Conflict of interest.** None declared.

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