Genome-Wide Prediction of the Polymorphic Ser Gene Family in *Tetrahymena thermophila* Based on Motif Analysis

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Abstract

Even though antigenic variation is employed among parasitic protozoa for host immune evasion, *Tetrahymena thermophila*, a free-living ciliate, can also change its surface protein antigens. These cysteine-rich glycosylphosphatidylinositol (GPI)-linked surface proteins are encoded by a family of polymorphic Ser genes. Despite the availability of *T. thermophila* genome, a comprehensive analysis of the Ser family is limited by its high degree of polymorphism. In order to overcome this problem, a new approach was adopted by searching for Ser candidates with common motif sequences, namely length-specific repetitive cysteine pattern and GPI anchor site. The candidate genes were phylogenetically compared with the previously identified Ser genes and classified into subtypes. Ser candidates were often found to be located as tandem arrays of the same subtypes on several chromosomal scaffolds. Certain Ser candidates located in the same chromosomal arrays were transcriptionally expressed at specific *T. thermophila* developmental stages. These Ser candidates selected by the motif analysis approach can form the foundation for a systematic identification of the entire Ser gene family, which will contribute to the understanding of their function and the basis of *T. thermophila* antigenic variation.

Introduction

*Tetrahymena thermophila* is a single-celled ciliate found in temperate freshwater [1,2]. *T. thermophila* naturally feeds on bacteria, but it can also grow in media under laboratory conditions [3]. It has two nuclei, a micronucleus (MIC) as a germ line and a macronucleus (MAC) as a source specific for gene expression [4]. It has two nuclei, a micronucleus (MIC) as a source specific for gene expression [4]. During the development of the new MAC, the parental MAC undergoes DNA rearrangements, including deletion of internal eliminated sequences (IES), removal of repetitive sequences and chromosome breakage at specific sites [4]. During the development of the new MAC, the parental MAC becomes degraded [4,5]. This unique biology of *T. thermophila* makes it an important model organism leading to seminal discoveries in the field of molecular biology [2].

*T. thermophila* cell membrane is covered by a surface protein known as immobilization antigen (i-ag) [6,7], as incubation with antibodies against i-ag causing *T. thermophila* to cease its movement, hence the name. Various subtypes of *T. thermophila* i-ag have been described based on immobilization assays with specific antibodies. Subtypes H, L and T are expressed at different temperatures [6,8], with subtype H being expressed at “high” temperature (20–35°C), subtype L at “low” temperature (<20°C) and subtype T (torrid) at temperatures above 36°C [8,9]. Gradual switching of i-ag subtypes occurs when temperatures are shifted [6,8]. The gene coding for i-ag was named Ser after the word “serotype” [10]. So far, up to six subtype H allelic variants, one subtype J gene and six subtype L paralogs were found [11–15]. One common characteristic among these Ser proteins is a repetitive cysteine-rich motif [13,14]. Such features are also common in surface proteins of other unicellular eukaryotes [16]. The control mechanism of Ser expression is not well understood, but the mRNA half-life of SerH3 (normally expressed at 20–35°C) is rapidly decreased when temperature is shifted up to 40°C [17,18]. Treatment with protein synthesis and protein kinase inhibitors can prolong SerH3 mRNA half-life during this temperature shift, suggesting that there are proteins and phospho-proteins involved in Ser mRNA degradation [19]. However, the role of *T. thermophila* i-ag remains unclear, though it may involve sensing the environment or prey-predator recognition, similar to *Paramecium* surface antigen [20].

Even though sequence variation is a hallmark of these highly diverged surface proteins, they often contain repetitive cysteine-rich motifs. The periodic cysteine residues could form disulfide bonds in a consistent pattern among proteins in the same family, but there is no experimental data on the formation of disulfide bonds in i-ag at this point. The existence of the disulfide bonds might introduce an extremely hydrophobic moiety at the core of...
protein [21]. It was suggested that 'hydrophobic collapse' might play a crucial role in protein folding in general via hydrophobic core nucleus which drives the folding process [22]. Disulfide bond formation could allow protein to become highly divergent on the surface while maintaining the overall fold. Different number of cysteine pair per one repeating sequence block in different i-ag subtypes has been documented [12]. Despite high sequence variation among Ser genes, the pattern of cysteine rich motif is a common feature among them.

Another distinct feature among Ser genes is the consensus sequence at the C-terminus specific for glycosylphosphatidylinositol (GPI) anchor modification. T. thermophila i-ag subtype H was shown to be GPI-anchored protein by radiolabelling [23,24]. Putative GPI anchor site was predicted to be located at the C-terminus of the Ser proteins [13,14]. There is a GPI anchor signal sequence at the C-terminus, which can be recognized by transamidase. GPI anchor signal can be divided into three regions: GPI attachment site, spacer region of polar residues (ω+3 to ω+8), and hydrophobic region (ω+9 to the C-terminus) [25]. There may be the minimum length for hydrophobic region required for GPI attachment [26]. Because GPI signal has no detectable conserved sequence, it cannot be identified by sequence similarity approach. However, the signal sequences surrounding the GPI attachment site can be defined as regions of amino acid residues with different physical properties such as size and hydrophobicity [25]. This allows the prediction of GPI-anchored protein using knowledge-based algorithm [27].

There are 24,725 predicted protein-coding genes in T. thermophila MAC genome [28,29]. The majority of the genes in T. thermophila are transcriptionally regulated as determined by nuclear run-on assay [30]. Post-transcriptional control via mRNA stability was also observed in SerH3 gene regulation [18]. As the Ser gene family is highly polymorphic, sequence homology analysis cannot recognize the full set of putative Ser genes.

In order to gain a full understanding of the antigenic variation in T. thermophila Ser genes, it is necessary to overcome this problem. In this study, the repertoire of the T. thermophila Ser gene family was annotated by setting search criteria based on the repetitive cysteine-rich motif and the signal sequence for GPI anchor. Two hundred and sixteen putative Ser genes from T. thermophila MAC genome sequence were selected including the known Ser genes. Previous studies have shown that each Ser subtype has a specific number of Cys residues per repetitive block: SerH with 8 Cys residues per block [14,15], SerJ with 10 Cys residues per block [12] and SerL with 6 Cys residues per block. The periodic cysteine block pattern, CXω+6Cω+6Xω+1Cω+9, and the GPI anchor site were used as the search criteria for the Ser gene family (Figure 1 and Figure S1 in File S1) [13–15]. Putative Ser genes were further characterized based on their phylogenetic distribution with experimentally classified Ser subtypes. By combining the data from the Ser candidates and the known Ser genes, the patterns of Ser chromosomal localization and gene expression were revealed.

Materials and Methods

Sequence and genome data
The whole predicted protein sequences of Tetrahymena thermophila were downloaded from Tetrahymena Genome Database (http://ciliate.org/index.php/home/downloads). The 5' UTR sequences of Ser candidates were obtained from TetraFGD Genome Browser (http://tdiglab.ibh.ac.cn). Thirteen known Ser subtypes (listed in Table S1 in File S1) from NCBI database were used in this study as reference [28,29].

Identification of Ser gene candidates
Ser prediction algorithm was composed of two main parts, cysteine pattern search and GPI-anchored protein prediction (Figure 2). Custom Perl script was used to perform the search for the cysteine-
rich pattern. The script was set to select any sequence containing $\text{CX}_{\leq 6}\text{CX}_{\leq 1}\text{CX}_{\leq 1}\text{CX}_{\geq 1}\text{CX}_{\geq 1}\text{CX}_{\geq 1}\text{CX}_{\geq 1}\text{CX}_{\geq 1}\text{CX}_{\geq 1}$ where match does not exceed 120aa within substring. This length of amino acid residues was chosen because it gave the best pattern match to the known Ser genes. The results were then refined by GPI-anchored protein prediction employing the web-based program FragAnchor [31]. Hidden Markov Model (HMM) imple-
Amino acid sequences matching the defined cysteine pattern with highly probable GPI-anchored protein determined by FragAnchor were selected as Ser candidates.

Phylogenetic analysis

The translated sequences of Ser candidates were aligned using ClustalX 2.0.12 with default multiple alignment parameters (gap opening penalty = 10; gap extension penalty = 0.2; Gonnet series weight matrix). Alignment was then adjusted manually. Neighbor-Joining (NJ) tree was calculated with 1000 bootstrap replicates using ClustalX. Protein evolutionary model was selected by ProtTest [32]. Maximum likelihood (ML) tree was estimated using RAxML 7.2.8 as implemented on the CIPRES Science Gateway [33–35]. Phylogenetic trees were then created using Dendroscope [36]. THERM_001980 was excluded from phylogenetic analysis because its sequence has an unusual sequence length of 3751 amino acid residues, preventing it from being aligned.

Figure 3. Neighbor-joining cladogram of Ser candidates identified in this study. Known Ser proteins are also included in this figure (black dots). 100-replicate bootstrapping was performed. Group-assigned candidates are highlighted. Bootstrap support percentages for the Neighbor-Joining (NJ) tree and the Maximum Likelihood (ML) tree (dash indicates undetermined bootstrap support value) are respectively shown on selected branch nodes. Among group-assigned candidates, THERM_00263370 and THERM_00329880 (marked with asterisk) are inconsistent between NJ and ML trees. In NJ tree, THERM_00263370 is clustered with L* clade. But in ML tree, it is closer to SerL clade. THERM_00329880 is clustered with SerH clade in the ML tree but not in the NJ tree.

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Ser gene expression analysis

*T. thermophila* genome-wide gene expression (growth, starvation and conjugation) microarray data was retrieved from *Tetrahymena* Functional Genomics Database (TetraFGD) (http://tfgd.ihb.ac.cn/) [37]. MultiExperimentViewer (MeV) was employed to evaluate gene expression clusters and to classify the

![Figure 4. Distribution of Ser candidates on T. thermophila MAC scaffold.](image)

Each short horizontal line represents one Ser gene. Ser subtype is color-coded. Dash line represents non-Ser gene which locates within Ser tandem arrays. Gene orientation on plus or minus strand is depicted by left or right alignment respectively. Scaffold number is shown on the top of each scaffold line (vertical line). Only scaffolds containing classified Ser tandem array are shown.

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expression pattern into subgroups (www.tm4.org/mev/). Clustering was identified with K-Means clustering module (KMC), and distance was calculated by Pearson correlation.

Results

Identification and classification of Ser genes

Due to the high degree of polymorphism, identification of T. thermophila Ser genes by sequence homology alone is limited by low sequence conservation. In order to systematically search for Ser candidates, two criteria were applied based on the common features found in existing i-ag proteins (six SerH, one SerJ and six SerL) namely, the presence of Cys residue pattern block CXlongCXshortC and the GPI-anchor signal located at the C-terminus. For the first criterion, a Perl script was set up to search for the T. thermophila proteins which, for any window frame containing 4 or 6 Cys residues within 30–120 amino acid residues, the number of amino acid between the first Cys pair is more than 5 and the number of amino acid between the other Cys pairs is at least 1 (Figure 2). After determining the number of hits versus saturation in gene numbers and change in phylogenetic pattern, the search criterion was limited to the minimum of 6 Cys residues within a 120 amino acid interval. Any protein containing CX($6$)CX($1$)CX($1$)CX($1$)CX($1$)C sequence within 120 amino acid residues was selected, resulting in 4,925 hits out of 24,725 T. thermophila predicted proteins. For the second selection criterion, FragAnchor classified 216 Ser candidates as “highly probable” GPI-anchored proteins including all known Ser genes (Figure 1–2 and Figure S1 in File S1). Sequences classified as “probable” or “weakly-probable” GPI-anchored proteins were excluded in order to avoid false positives. The approach successfully identified all four annotated Ser genes in the genome of T. thermophila strain SB210 with perfect identity match to the experimentally identified Ser genes.

Classification of putative Ser genes from T. thermophila strain SB210

Forty-five percent of the selected genes could be grouped with three identified Ser subtypes, SerH, SerL and SerJ, with good bootstrap support from NJ analyses (SerH: 58.7% NJ bootstrap support; SerL: 72.8% NJ bootstrap support; and SerJ: 99.8% NJ bootstrap support). Two distinct branches of Ser candidates, grouped with SerJ and SerL, were found and were named J* (74.6% NJ bootstrap support) and L* (42.2% NJ bootstrap support) to reflect their phylogenetic association with SerJ and SerL groups, respectively (Figure 3). Candidates grouped with the known Ser subtypes are listed in Table S2 in File S1.

Ser chromosomal location

Chromosomal location analysis focused on the newly identified Ser genes with known subtypes. They were dispersed on 76 out of the estimated 250–300 MAC chromosomes in T. thermophila. Figure 4 shows the distribution of subtype-classified Ser candidates on 12 scaffolds, with the majority (86%) of the identified Ser candidate genes located in close proximity to one another. Ser genes tend to form a tandem array of the same subtype (Table S2 in File S1). Six MAC scaffolds contained tandem arrays with only

![Figure 5. Ser gene expression clusters.](image)

Each row represents one gene. Gene ID and subtype were listed. Unclassified Ser candidates are marked as X. Gene expression data was subjected to K-means clustering method using Pearson correlation to measure distance. Expression cluster ID is indicated as arabic number on the left of expression heatmap. Scale bar represents log2-transformed gene expression value. Red indicates expression value above median, and green indicates expression value below median. Clustering analysis was performed using MeV 4.7. doi:10.1371/journal.pone.0105201.g005

![Figure 6. Expression profile of Ser genes from tandem array 38-1 and 60-1.](image)

Gene expression data was collected from T. thermophila during growth at low (Ll), medium (Lm), and high (Lh) densities. The data from starvation (S0, S3, S6, S9, S12, S15, S24) and conjugation (C0, C2, C4, C6, C8, C10, C12, C14, C16, C18) samples was also included with numerical values showing hours in each particular condition [37]. Complete Ser expression cluster result is available in Table S2 and S3 in File S1. doi:10.1371/journal.pone.0105201.g006
one subtype of Ser genes, and those in close proximity on the same scaffold tended to have the same orientation, with a few exceptions, such as subtype-H TTHERM_00602920 on scaffold 84 and subtype-L TTHERM_0099320 on scaffold 3835. There appears to be no preference for scaffold size or chromosomal region where Ser genes are located.

Ser gene expression analysis

Ser expression patterns during growth, starvation and conjugation were analyzed based on DNA microarray data [37]. Ser expression data could be grouped into 30 clusters based on their expression patterns (Figure 5). No correlation was found between specific Ser subtypes and expression patterns. Certain Ser tandem genes were associated with the same expression pattern. For example, ten out of thirteen subtype-L Ser genes on scaffold 38 were found in expression cluster 4 (Figure 6, upper panel). However, some tandem arrays were not in the same expression cluster, but their expression patterns appeared to be stage-specific. For example, scaffold 60 contains a tandem array of 6 subtype-H Ser genes grouped into four different expression clusters (Figure 6, lower panel). Interestingly, they were all up-regulated during conjugation, but at different time points. Data of similar analyses on locations and expression patterns of unclassified Ser candidates are available in Table S3 in File S1.

Discussion

T. thermophila i-ag was originally identified as variant surface antigen. Their high degree of polymorphism has prevented systematic identification of its Ser family based on sequence homology alone despite available genomic data. In this study, an alternative strategy was adopted for identifying this gene family by selecting two common features of the known Ser genes, namely, the Cys-rich motif and GPI-anchor site. Their phylogenetic distributions were analyzed in order to characterize Ser candidates that are related to the known subtypes. The approach identified Ser candidates that can be grouped into the known subtypes. However, 55% of the genes could not be classified into any known subtype based on phylogenetic analysis, and might belong to new subtypes. At present, the genes encoding several types of surface antigens such as SerT, SerS, SerM and SerI have not been identified indicating the existence of more Ser repertoires and subtypes [30]. In addition, certain unclassified gene candidates exhibit properties similar to Ser, such as tendency to be located in tandem and have similar gene expression profile. It is possible that the missing Ser subtypes might belong to one of the unclassified gene families. Experiments with specific antibodies to Ser candidates will be required in order to prove that they are indeed i-ag proteins. Sequence comparison and syntenic analysis of the highly polymorphic Ser genes from more T. thermophila isolates will confirm whether these genes are under positive selection which is a strong evolutionary driving force for mating proteins, molecular sensors and evasive decoys [39,40]. Using the Ser candidates as blast query on NCBI protein database yields sixteen T. thermophila proteins not previously included in the list [Table S4 in File S1]. Nevertheless, they either lack the GPI anchor motif or matching cysteine pattern. The phylogenetic analysis showed that these proteins are not grouped with any Ser candidate. The analysis outside T. thermophila revealed two weak hits with the proteins from other ciliates [Ichthyophthirius multifilis AAK94941 with 21% identity to SerL and Plasmadium tetraurelia XP_001459224 with 22% identity to SerH]. This might indicate that the Ser gene family is unique to T. thermophila.

i-ag proteins were discovered based on their variation in response to antibodies directed at T. thermophila surface antigens. This is a hallmark for many antigenic variation phenomena among parasitic and free-living protozoa [41]. In parasitic protozoa such as Plasmadium falciparum, a family of proteins on infected red blood cells is needed for a parasitic adherence mechanism to human cells and tissue which is crucial for malaria pathogenesis [42]. Antigenic variation in P. falciparum switches the variant pathogenic proteins in order to avoid immune detection [43]. Other parasitic protozoa also exploit a similar system [44,45]. When antigenic variation is compromised, the parasite becomes vulnerable to the immune system [46].

Our study took advantage of the available T. thermophila MAC genome and DNA microarray expression data to analyze the Ser gene family. These Ser genes are organized in tandem arrays on several MAC scaffolds. These tandem arrays often belong to the same subtype, suggesting that they arose by gene duplication or genetic recombination. Expression pattern cluster analysis does not explicitly indicate the role of the Ser gene family in any developmental stage in particular. Available microarray data also did not include every known culture conditions for inducing the expression of Ser genes. However, stage-specific expression patterns of several Ser transcripts at the same time point were observed, and thus mutual exclusion mechanism is not likely the only strategy underlying expression control of every Ser gene.

Unlike parasitic protozoa, free-living protozoa are not subject to host immune pressure, and the purpose for having surface antigenic variation remains unclear. Nevertheless, understanding the mechanism of antigenic variation in free-living organisms could provide new insights into the evolution and regulation control of antigenic variation in parasitic organisms. Thus identifying the whole repertoire of the Ser gene family is the first step towards the exploration of antigenic variation in T. thermophila, an important model organism for many seminal discoveries in molecular biology.

Supporting Information

File S1 Supporting information. (DOC)

Acknowledgments

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Author Contributions

Conceived and designed the experiments: PP TC. Performed the experiments: PP. Analyzed the data: PP KK TC. Contributed reagents/materials/analysis tools: PP. Wrote the paper: PP KK TC.

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