Functional Analysis of Hypothetical Proteins of *Chlamydia Trachomatis*: A Bioinformatics Based Approach for Prioritizing the Targets

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Abstract

The various genome sequencing projects have led to the accumulation of entire set of gene sequences of many organisms. Among the sequenced genomes are numerous genes which code for proteins of unknown function. These genes are termed as hypothetical genes and their corresponding gene products are known as Hypothetical Proteins (HPs). Analyzing and annotating the functions of these HPs is important in pathogenic organisms such as *Chlamydia trachomatis* that causes various sequelae of diseases by infecting different sites in humans. Functional annotations of these HPs provides insights into their exact molecular function and may help in identification of novel drug or vaccine candidates for the control of infections caused by *C. trachomatis*. In the present study, entire set of 336 HPs of *C. trachomatis* were retrieved from NCBI and analyzed for their function using bioinformatics tools such as CDD-BLAST, PFAM, TIGRFAM and SCANPROSITE. The analysis revealed that some of the HPs corresponded to the bacterial secretory pathway proteins. The structural prediction of the annotated proteins has been performed which further substantiate the functional characterization results. Bioinformatics approach used in this study, including sophisticated sequence analysis, domain characterization and structural prediction studies, can provide a useful lead to experimentally annotate and corroborate these studies. Data generated by this study might facilitate swift identification of potential therapeutic targets and thereby enabling the search for new inhibitors or vaccines.

Keywords: *Chlamydia trachomatis*; Hypothetical proteins; Drug targets; Vaccine candidates

Introduction

*Chlamydia trachomatis* infection is one of the most common sexually transmitted diseases worldwide, with an estimated 7.2 million disease cases annually, of which majority of the cases occur in Asia (including India), Sub-Saharan Africa and South America [1-3]. Moreover, in India we have limited capacity to effectively screen for infections caused by *C. trachomatis* and only few labs are actively working in this area [4-9]. Since it is well established that chlamydial infections can lead to a variety of asymptomatic and symptomatic manifestations including vaginal mucous-purulent discharge, endometritis, salpingitis and Pelvic Inflammatory Disease (PID) [10], it becomes imperative to screen for such infections at an early stage. In women, the infection is mostly asymptomatic and among the infected women, it is estimated that approximately 20% develop PID, 4% chronic pelvic pain, 3% become infertile and 2% have an adverse pregnancy outcome [11-13]. These complications increase further because of the occurrence of numerous *C. trachomatis* serovars, of which eight serovars (D, E, F, G, H, I, J, K) are known to cause genital infections [14].

In view of the observed complications, it is clear that not only screening programs are important, proper treatment regimens should also be followed for its effective control. In the pre-antibiotic era, chlamydial infections were often life threatening and even today they may give rise to death despite treatment with antibiotics as it has been established that some of the *C. trachomatis* serovars can produce a number of different components that may contribute to virulence, eventually leading to death of a patient [10]. Although, antibiotic therapy is thought to eliminate these chlamydial infections [15] and thereby prevent such extreme cases, it does not treat the established pathology. This, together with the fact that chlamydial infections can often be asymptomatic, also point towards the urgent need for the development of preventative measures such as vaccination for control of the disease. Moreover, as per the guidelines of World Health Organization (WHO), treatment is largely based on symptomatic case management [1]. Since a large proportion of infective population remains asymptomatic, treatment on the basis of such guidelines not only misses out on asymptomatic patients but could also lead to over treatment. It is important to understand that treatment regimen should be based on local antibiotic sensitivities but such pattern is rarely known in resource limited settings such as India and hence may contribute to emergence of resistant strains of this pathogen which may further spread to other countries as well. Even though, today most of the antibiotics are effective against *C. trachomatis* infections but the time is not far wherein the antibiotics would turn out to be ineffective as with other infectious diseases. Therefore, it is becoming increasingly important that we not only explore novel drugs / targets for its treatment but also develop new vaccine candidates as a step towards prevention of the disease. Unfortunately, development of novel anti-chlamydial drugs is hampered because of our lack of understanding of its pathogenesis. Moreover, *C. trachomatis* infection can be prevented...
worldwide, once we are able to develop newer vaccines as current vaccines are no longer effective against Chlamydial infections owing to the presence of multiple serovars. These two objectives, i.e., novel drug targets and/or vaccine development could be readily achieved if one is able to understand Chlamydial biology in a holistic manner. Along with experimental studies, bioinformatics and genomics might play an important role in achieving the above mentioned objectives, as they have done for other infectious organisms. Over the last decade, more than 150 complete genomes of diverse bacteria, archaea and eukaryotes have been sequenced and many more are currently in pipeline [16,17]. Although, a large number of microbes have been sequenced, about 50% of genes do not show homology to functionally characterized proteins. Functional annotation of these HPs might offer an opportunity to understand novel drug designing methods and vaccine development.

In the present study, we have made an attempt to exploit the data generated using bioinformatics tools to help us investigate newer treatment options/preventive measures available for the pathogenic organism, C. trachomatis. The completed genome sequence for C. trachomatis D/UW-3/CX provided us with a comprehensive inventory of all the proteins potentially produced by this organism. A significant question that we wanted to answer was as to how this information can guide us in identifying novel drug targets or vaccine candidates. The Chlamydia whole genome is a single chromosome which contains 1042519 base pairs (bps) [-1.04 megabase pairs (MB)] having Adenine and Thymine (AT) content of 58.7%. Total number of protein encoding genes throughout the C. trachomatis genome is 894 (genome sequencing and annotation methodologies available are at the Science website: (www.sciencemag.org/feature/data/982604/.shl)). Out of these 894 protein coding genes, 558 (62%) genes encode for proteins that have assigned function and 336 (38%) genes are hypothetical in nature.

Hypothetical Protein (HP) is a protein that is predicted to be expressed from an open reading frame, but for which there is no experimental evidence of translation. These constitute a substantial fraction of proteomes of prokaryotes and are even present in eukaryotes [16]. With the general belief that the majority of HPs are the product of pseudogenes, it is essential to have a tool with the ability of pinpointing the minority of HPs with a high probability of being expressed. In the strict sense, HPs are predicted proteins, as predicted from their nucleic acid sequences and that have not been shown to be expressed by experimental procedures. Moreover, these proteins are characterized by low identity to known, annotated proteins. Among these HPs, a separate class of "conserved hypothetical" proteins are defined as a large fraction of genes in sequenced genomes encoding those that are found in organisms from several phylogenetic lines but have not been functionally characterized and described at the protein chemical level [18]. These structures may represent up to half of the potential protein coding regions of a genome.

In order to treat infectious diseases such as those caused by C. trachomatis, functional annotation of these HPs might open avenues for prioritizing vaccine candidate genes or novel drug targets. Structural genomics initiatives provide ample structures of hypothetical proteins at an ever increasing rate. However without functional annotation of these proteins, this structural analysis would be of no use to biologists who are always interested in deciphering particular molecular mechanisms. Moreover, some of the proteins, which are considered to be well annotated, may have additional functions beyond their listed records. Undertaking such studies, a series of additional protein pathways and cascades can be revealed, completing our fragmentary knowledge on the mosaic of proteins per se. Lastly we may emphasize that the analysis of HPs would be of benefit to genomics enabling the discovery of so far unknown or even predicted genes. These annotated HPs may serve as markers and pharmacological targets in the era of personalized medicine for C. trachomatis.

Methods

Sequence retrieval and functional annotation

Complete genome sequence of C. trachomatis D/UW-3/CX was retrieved from NCBI database (Sequence and annotation available at (http://chlamydia-www.berkeley.edu) and GenBank under accession AE001273) and the sequences of "hypothetical proteins" of Chlamydia trachomatis were analyzed. Out of 894 protein sequences, 336 hypothetical protein sequences were analyzed for the presence of conserved domains using sequence similarity search with orthologous family members available in various databases using web-tools.

Four bioinformatics tools CDD-BLAST (http://www.ncbi.nlm.nih.gov/BLAST/) [19-21], TIGRFAM (http://www.jcvi.org/cgi-bin/tigrfams/index.cgi) [22], PFAM (http://www.pfam.sanger.ac.uk) [23] and SCANPROSITE (http://prosite.expasy.org/scanprosite) [24] were used, which can search the defined conserved domains in the targeted protein sequences and further assist in the classification of putative proteins in a particular protein family.

CDD-BLAST is NCBI’s web interface to search the Conserved Domain Database with protein query sequences. It uses RPSBLAST, a variant of PSI-BLAST, to quickly scan a set of pre calculated Position-Specific Scoring Matrices (PSSMs) with a protein query. PFAM is a collection of multiple protein-sequence alignments and Hidden Markov Models (HMMs) and provides a good repository of models for identifying protein families, domains and repeats. TIGRFAM is a manually curated database which exploits HMMs to annotate protein families by search alignment for each family. It contains more than 1600 protein families and gives a base through which one can extract a number of information regarding genome annotation, sequence similarity, sequence alignment, function profiling and phylogenetics profile analysis during model construction. SCANPROSITE is an improved version of web based tool which detect PROSITE signature matches in given protein sequence.

Functional categorization

Hypothetical proteins analyzed by the above mentioned function prediction web tools have shown the variable results when searched for the conserved domains in hypothetical sequences and different confidence levels have been generated on the basis of collective results of these web-tools.

1. If the given four tools indicate the same functions then the confidence level were to be 100 percent.
2. If the given three tools indicate the same functions and one is showing different function then the confidence level were to be 75 percent.
3. If the given two tools indicate the same functions and two are showing different functions then the confidence level were to be 50 percent.
4. If only one tool indicates the function and other tools are showing different functions then the confidence level were to be 25 percent.
Protein structure prediction

Predicting the 3D structure of a HP from its amino acid sequence using computational methods also allows us to predict its function to an extent [25]. Two types of tertiary structure prediction methods are known namely, ab initio methods which predict a protein structure based on physico-chemical principles directly, and template-based methods, which uses known protein structures as templates. Template based methods include homology or comparative modeling, and fold recognition via threading.

In our study we have used an online server PS2-v2 (PS Square version 2) Protein Structure Prediction Server [26] which is a template based method to predict the structure of HPs. PS2-v2 uses the strategies of Pair-wise and multiple alignments by combining powers of the programs PSI-BLAST, IMPALA and T-COFFEE in both target – template selection and target–template alignment and finally it constructs the protein 3D structures using integrated modeling package of PS2 using best scored orthologous template.

We found that (PS2-v2) server to be a very user-friendly web server. The query protein sequence of the HP was given as an input in FASTA format. The server provided three modes (Automatic, Manual and 'Use this template') for choosing the template. The default mode was 'Automatic'. In the automatic mode, (PS2-v2) automatically selects the modeling template(s). For the ‘Manual’ mode, server enables users to assign specific template(s) from a list of candidates. The ‘Use this template’ mode allows users to assign a specific protein structure as the template. We used the default mode for all the eleven proteins in order to predict their structures. Predicted results were received by us later at our email address. Output of the server showed a list of templates, selected template(s), target-template alignment(s), predicted structure(s) and structure evaluations as the result.

Results and Discussion

Advances in high throughput modern DNA sequencing technologies combined with its cost efficiency has enabled sequencing of a large number of bacterial genomes. Since many of the genes are conserved across a large number of bacterial genomes, accurate annotation of the genes usually relies on sequence homology methods wherein, the function of a specific gene is assigned based on sequence similarity to gene with known function [27]. In spite of this enormous proliferation of genomic data more than one third of genes have no assigned function. One of the reasons is the functional divergence of similar sequences during the course of evolution [28,29]. Hence sequence homology based methods alone fail to assign accurate functions to a large number of genes and may lead to imprecise annotations [30]. To circumvent this problem, multiple tools should be used to assign functions to hypothetical proteins, as this may help in reducing the fraction of HPs considerably. The present study concentrated on the functional annotation of hypothetical proteins from C. trachomatis using diverse bioinformatics tools. The four webtools used in the current study helped us to search the presence of conserved domains in 336 Hypothetical Proteins (HP). Domains are often identified as recurring (sequence or structure) units, which may exist in various contexts and can be thought of as distinct functional and/or structural units of a protein. In molecular evolution such domains may have been utilized as building blocks and may recombine in different arrangements to modulate protein function [31]. Once the presence of a specific domain in a protein was established, we further classified proteins into various categories. Depending on our results, as represented in Table S1, we found that there were 11 proteins having a consensus in the presence of the domain using all the four bioinformatics tools and hence they were grouped together under the confidence limit of 100%, as shown in Table 1.

Out of the remaining 325 proteins, we could not find any specific domains for 83 proteins using the four bioinformatics tools. For these 83 proteins, structural analysis might provide some meaningful results. For other hypothetical proteins (n=242), specific domains were identified using one, two or three of the above mentioned tools. Accordingly these were categorized under the confidence limit of 25% for 134 proteins, 50% for 66 proteins and 75% for 42 proteins (Table 1). These may or may not be specific for the highlighted domains. The exact function of these proteins requires further studies.

Within a protein family, a domain or fold may be more strongly conserved than the entire sequence [32]. Amongst the 11 HPs for which functional domain was identified with a confidence level of 100%, three proteins (CT110, CT341, CT396) were classified as chaperonins (Table 2). Other proteins showed domains suggestive of their function as ligase, synthase, protease, bacterial secretory pathway proteins, translocase and a zinc finger domain carrying protein suggesting that these proteins of Chlamydia may be involved in performing similar function. Presence of zinc finger domain in CT407 implies that it might be a DNA binding protein, probably a transcription regulator. The classification of the HPs according to the presence of specific domains and their super-family descriptions is given in Table 2.

Biochemical function of the protein can also be verified by inclusion of structural information [33]. The three dimensional structures of the eleven annotated proteins were modeled by PS square (PS2-v2) online server, which is dependent on Template-Based Modeling (TBM) and fold recognition methods. Using this method, a prediction model is built based on the coordinates of the appropriate template of the protein. These approaches generally involve four steps: 1) a representative protein structure database is searched to identify a template that is structurally similar to the protein target; 2) an alignment between the target and the template is generated that should align equivalent residues together as in the case of a structural alignment; 3) a prediction structure of the target is built based on the alignment and the selected template structure, and 4) model quality evaluation. The first two steps significantly affect the quality of the final model prediction in TBM methods. The templates used by the server to model these proteins are tabulated in Table 3. The modeling of CT110, CT341, CT396 proteins using PS square (PS2-v2) online server further substantiated the chaperonin function of these three proteins.

### Table 1: Confidence level of All Four Tools Used in this Study (Total proteins 336).

<table>
<thead>
<tr>
<th>Percentage of similarity</th>
<th>0%</th>
<th>25%</th>
<th>50%</th>
<th>75%</th>
<th>100%</th>
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<tbody>
<tr>
<td>Number of hypothetical proteins (Total 336)</td>
<td>83</td>
<td>134</td>
<td>66</td>
<td>42</td>
<td>11</td>
</tr>
</tbody>
</table>

### Table 2: Putative Functional Category of Hypothetical Proteins with 100% confidence level According to the Functional Domain They Contain (Total 11 Proteins).

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Functional categories to which they belong</th>
<th>List of proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Chaperonin</td>
<td>CT110, CT341, CT396</td>
</tr>
<tr>
<td>2.</td>
<td>Ligase</td>
<td>CT146</td>
</tr>
<tr>
<td>3.</td>
<td>Synthase</td>
<td>CT171, CT257</td>
</tr>
<tr>
<td>4.</td>
<td>Protease</td>
<td>CT431, CT706</td>
</tr>
<tr>
<td>5.</td>
<td>Bacterial secretory pathway proteins</td>
<td>CT571</td>
</tr>
<tr>
<td>6.</td>
<td>Translocase</td>
<td>CT701</td>
</tr>
<tr>
<td>7.</td>
<td>Zinc finger protein</td>
<td>CT407</td>
</tr>
</tbody>
</table>
found to be functionally significant. In comparison to their approach, which were Haemophilus influenzae, Streptococcus pneumoniae and Haemophilus parainfluenzae, the group was able to identify 8 potent target genes from a gene based on synonymous codon usage, is coupled with subtractive strategy. The annotation allows scientists to reduce the number of uncharacterized HPs, which produce many virulence factors and cause serious infections and disease complications, is still ambiguous. Further understanding of functional properties of these HPs of C. trachomatis will not only provide a better insight into the pathogenesis of C. trachomatis, but may also help in identifying novel therapeutic candidates. In fact, there has been a recent report on genome evolution. They predict functional associations between protein coding genes by analyzing gene fusion events, the conservation of gene neighborhood, or the significant co-occurrence of genes across different species. Unlike homology based annotation, which infers molecular features by information transfer from experimentally characterized proteins, genomic context methods predict functional associations. Therefore, it should be noted that these methods do not provide information about exact biochemical or experimental function and are time consuming. In silico analysis described here provide an easy and accurate method of assigning function for various HPs. Nevertheless using systematic hierarchical approaches together, would help scientists in reducing the number of uncharacterized HPs.

Although we have been able to accomplish a comprehensive in silico analysis, still function of around 300 genes of the studied pathogenic bacteria C. trachomatis, which produce many virulence factors and cause serious infections and disease complications, is still ambiguous. Further understanding of functional properties of these HPs of C. trachomatis will not only provide a better insight into the pathogenesis of C. trachomatis but may also help in identifying novel therapeutic candidates. In fact, there has been a recent report wherein a novel genomics approach using Codon Adaptation Index (CAI), a measure that is used to predict the translational efficiency of a gene based on synonymous codon usage, is coupled with subtractive genomics approach for mining potential drug targets. Using the strategy, the group was able to identify 8 potent target genes from Streptococcus pneumoniae and Haemophilus influenzae, which were found to be functionally significant. In comparison to their approach, our study facilitates swift identification of the hidden function of HPs which could become potential therapeutic targets and may indicate a key role in host-pathogen interactions. Once established as novel drug or vaccine targets, further research for new inhibitors and vaccines can be accomplished.

### Conclusions

In the present study, we have taken a close look at 336 chlamydial HPs whose function was analyzed using diverse publicly available bioinformatics tools. Based on the various domains studies, we were able to classify 11 hypothetical proteins under different functional categories. The analysis revealed functionally important domains and families which were involved in inducing protein synthesis and multiple antibiotic resistances in the bacteria and also perform enzymatic functions. A few of the annotated proteins could turn out to be novel targets to combat antibiotic resistance and vaccine development once their role is validated experimentally. Future studies may be directed towards subcellular localization of these annotated proteins for elucidating their cellular processes. Knowledge of the subcellular localization of a protein can significantly improve target identification during the drug discovery process. Once the structures of these annotated proteins is established and their function is known, further investigation into their ligand binding sites would help us identify newer antimicrobials against resistant strains. Currently there is also an immediate need for newer and effective vaccine providing protection against Chlamydia trachomatis due its inherent antigenic variation. Proteins that are secreted by various microbes to the extracellular environment could turn out to be useful antigens that might induce protective immunity or could elicit an immune response of diagnostic value. Above mentioned subcellular localization studies would be, therefore, beneficial in identifying such antigenic proteins. Finally we may emphasize that quantitative computational analysis, carried out in the present study, may help us in better understanding of the biology of Chlamydia trachomatis as a whole and identify potential therapeutic leads at the molecular level.

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### References


<table>
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<th>Structure</th>
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</tr>
<tr>
<td>2.</td>
<td>CT146</td>
<td>2w0A</td>
<td>Ligase</td>
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<td>3.</td>
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<td>2kxB</td>
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<td>4.</td>
<td>CT257</td>
<td>2yyA</td>
<td>Transporter</td>
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<tr>
<td>5.</td>
<td>CT341</td>
<td>1n1A</td>
<td>Chaperonin</td>
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<td>6.</td>
<td>CT396</td>
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</tr>
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<td>7.</td>
<td>CT407</td>
<td>1jGA</td>
<td>Zinc Finger Protein</td>
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<td>CT431</td>
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<td>Protease</td>
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</tr>
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</tr>
<tr>
<td>11.</td>
<td>CT706</td>
<td>1y6A</td>
<td>Protease</td>
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</tbody>
</table>

**Table 3:** Structural prediction of annotated proteins using the (PS)2-v2: template-based protein structure prediction server.


