**UTMGO: A Tool for Searching a Group of Semantically Related Gene Ontology Terms and Application to Annotation of Anonymous Protein Sequence**

Razib M. Othman, Safaai Deris, and Rosli M. Illias

**Abstract**—Gene Ontology terms have been actively used to annotate various protein sets. SWISS-PROT, TrEMBL, and InterPro are protein databases that are annotated according to the Gene Ontology terms. However, direct implementation of the Gene Ontology terms for annotation of anonymous protein sequences is not easy, especially for species not commonly represented in biological databases. UTMGO is developed as a tool that allows the user to quickly and easily search for a group of semantically related Gene Ontology terms. The applicability of the UTMGO is demonstrated by applying it to annotation of anonymous protein sequence. The extended UTMGO uses the Gene Ontology terms together with protein sequences associated with the terms to perform the annotation task. GOPET, GOtcha, GoFigure, and JAFa are used to compare the performance of the extended UTMGO.

**Keywords**—Anonymous protein sequence, Gene Ontology, Protein sequence annotation, Protein sequence alignment

**I. INTRODUCTION**

The Gene Ontology (GO) [1] is a project to provide a rich and comprehensive unified vocabulary to describe genes and their functions and products. The vocabulary is formed as a hierarchy of terms in three main categories: molecular function, biological process, and cellular component. Currently the GO comprises more than 20 thousand terms and is updated every 30 minutes, which tally with the growth activities in the bioinformatics field. The GO terms have been widely used for annotating various protein sets such as in DRTF [2], a database of rice transcription factor; SCOPPI [3], a database of protein domain-domain interactions; NOPdb [4], a database of nucleolar proteome; and Organelle DB [5], a database of protein localization and function.

One of the advantages of the GO terms is that it can cope with synonyms and can describe biological function. Furthermore, the GO terms are linked with approximately 8.2 million associations, 1.9 million different gene products, and with the largest set covering around 1.8 million protein sequences from 0.3 million species. Thence, specific protein sets can easily be compared with respect to common functional features [6], [7], protein databases such as MiGenes [8] and PA-GOSUB [9] can be explored through complicated queries, and large-scale protein database can simply be annotated [10], [11] based on the GO terms. However, direct use of the GO terms to annotate anonymous protein sequences is not easy, especially from small sequencing projects or for species not commonly represented in biological databases. Furthermore, for small group of scientists with little computational background or without appropriate facilities it is a tedious task to annotate those protein sequences.

In this paper, we present UTMGO, a tool for searching a group of semantically related GO terms. The basic structure of the UTMGO is extended to show how it could be applicable to annotation of anonymous protein sequence. Generally, the UTMGO consists of two primary components. The first component is applied to cluster the monolithic GO RDF/XML file into a set of more accessible and understandable files. The second component is employed to search for semantically related GO terms together with protein sequences associated with the terms from the fragmented GO RDF/XML files. The extended version of the UTMGO integrates JAligner engine [12] to perform protein sequence alignment. The JAligner engine has been modified to comply with the extended UTMGO. JAligner is a Java implementation of protein pairwise sequence alignment algorithms. This tool applies the dynamic programming algorithm Smith-Waterman. Both of the versions are described in detail in Section III.

This paper is organized as follows. Section II presents existing tools for annotating anonymous protein sequence. Section III gives the structure of the UTMGO and its extended version. Section IV demonstrates the working of UTMGO.

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II. EXISTING PROTEIN SEQUENCE ANNOTATION TOOLS

Annotation is a process of associating additional information with a particular point in a piece of information. Instead of annotating genomes and protein sets, it has been widely applied in annotating image [13] and web [14]. In the post-genomic era, annotation of protein sequence should be inferred from annotations of the nucleotide sequence, analogies with already understood proteins, plus references to patterns and motifs characteristic of particular protein functions. This can be achieved using the GO terms that have associations with gene products provided by Gene Ontology Annotation (GOA), see example in Fig. 1. In the GOA project which can be accessed at www.ebi.ac.uk/GOA/, electronic mappings and manual curation are used to assign the GO terms with all proteomes that exist in the UniProt Resource (UniProtKB/Swiss-Prot, UniProtKB/TrEMBL, and PIR-PSD), Ensembl, and other biological databases. The GOA statistics for Homo sapiens and other species are shown in Fig. 2.

A number of tools have been developed for annotation of anonymous protein sequence based on the GO terms. These

(both basic and extended versions) and its comparison with other tools. Section V concludes with a discussion and a brief outline of future improvement directions.

Fig. 1 Example of GO term. (A) The GO term, “DNA binding” (GO:0003677). (B) Gene ac (achaete), a gene product associated to “DNA binding” based on GOA. (C) Protein sequence of gene ac in FASTA format. (D) Synopsis of gene ac in FlyBase database. (E) Reference that relate “DNA binding” and gene ac in accordance with Traceable Author Statement (TAS) evidence.
GeneAtlas [27], and SMART [28]. There are also works using such as QuasiMotiFinder [24], MyHits [25], Pfaat [26], group of similar tools to annotate the anonymous protein computational linguistic techniques. On the other hand, a Verspoor 5) tools include:

1) GOPET [15] is an automated annotation tool for assigning the GO terms to cDNA or protein query sequences. It utilizes the GO for annotation terms, BLAST (Basic Local Alignment Search Tool) and GO-mapped protein databases for performing homology searches, and support vector machines for the prediction and the assignment of confidence values.

2) GOtcha [16] is a tool that provides a prediction of a set of GO terms for a given query sequence (DNA or protein). BLAST is used to get the initial score of each GO term and the scores calibrated against term-specific probability (P-score) to give higher accuracy.

3) GObolt [17] is a tool that offers annotation for anonymous cDNA or protein sequences according to the GO terms. It uses the GOA together with a series of protein databases and then employs BLAST to perform annotation by sequence similarity searches.

4) GoFigure [18] is a tool that accepts unknown DNA or protein sequence as an input and then uses BLAST to predict the GO terms by identifying homologous sequences in the GO annotated databases.

5) JAFA [19] is a meta-server that uses several function prediction programs such as GOtcha, GObolt, GoFigure, InterProScan [20], and Phydabac [21]. It accepts a protein sequence and provides predictions based on the GO terms.

In the meantime, an extensive study has been done by Verspoor et al. [22] and Xie et al. [23] counting on computational linguistic techniques. On the other hand, a group of similar tools to annotate the anonymous protein sequence without relying on the GO terms are also available such as QuasiMotifFinder [24], MyHits [25], Pfaat [26], GeneAtlas [27], and SMART [28]. There are also works using Bayesian method, statistical method, and C4.5 that have been carried out by [29]–[31] respectively.

However, little effort has been done to develop a tool with the following features to overcome the existing weaknesses:

1) Not dependent on BLAST, requires low cost and minimum hardware specifications, and with reasonable amount of execution time.

2) Not dependent on FASTA (Fast Alignment) format, the query protein sequence can be a newly sequenced one and not necessarily represented in existing biological databases.

3) Not dependent on RDBMS (Relational Database Management Systems), does not require user to setup the RDBMS software and to import the data or sources into the RDBMS format.

4) Fully based on the GO data without requiring download of GOA data or sequence sets from various sources.

5) Annotation is not only dependent on molecular function terms but also on biological process and cellular component terms.

6) Sequence alignment is not carried out to all protein sequences but only to sequences with higher outguessed similarity.

III. THE STRUCTURE OF THE BASIC AND EXTENDED UTMGO

The heart of the UTMGO is its searching and clustering engines, namely SSMGA [32] and SMAGA [33] respectively as shown in Fig. 3. Brief explanations of processing behind the UTMGO are as follows:

1) Public GO data in MySQL and RDF/XML formats are downloaded from the GO website.

2) The single GO RDF/XML file is split into smaller files by SMAGA. The number of files created and the size and the GO terms delegated to each file will be determined automatically during the execution of the clustering process.

3) Corresponding gene products together with protein sequences associated with the GO terms either based on IEA (Inferred from Electronic Annotation) or non-IEA evidence from the GO MySQL database are added into the fragmented GO RDF/XML files.

4) The UTMGO requires the user to enter a GO term and the number of matched GO terms to be returned $N_{gt}$.

5) Results which are $N_{gt}$ number of GO terms with higher term similarity score to the query GO term will be generated by SSMGA.

The UTMGO is a Linux based tool. Its engines are developed using C/C++ languages and it uses JSP (Java Server Pages) scripts for submitting queries via a web form. The extended version of the UTMGO, as shown in Fig. 4, is specifically designed for annotation of anonymous protein sequence. This extended version comprises of the following steps:

1) Get an anonymous protein sequence, the number of GO terms to be returned $N_{gt}$, a term similarity threshold, a number of protein sequences associated with each GO
term to be returned $N_t$, and optionally a GO term from the user.

2) If the GO term is void, then go to step 3. Otherwise, go to step 6.

3) Get input from the user for appropriate species, matrix type either BLOSUM (Blocks Substitution Matrix) or PAM (Point Accepted Mutations), and open and extend gap penalties to limit the search.

4) Perform sequence similarity search using the JAligner engine for the given anonymous protein sequence from step 1. The search is executed for protein sequences from the fragmented GO RDF/XML files that relate to the molecular function terms. The output will be a protein sequence with the highest sequence alignment score.

5) Select molecular function term with maximum cardinal of association with the protein sequence obtained from step 4 for the next step. If there is more than one term, the user will make the selection.

6) Submit the GO term either from step 1 or step 5 to the UTMGO and then perform term similarity search.

7) Return $N_t$ number of GO terms with term similarity score higher than the term similarity threshold from step 1.
thus, together with protein sequences associated with them.

8) Compute sequence alignment score between the query sequence and all sequences for each GO term gained from previous step using the JAligner engine. Display only $N_s$ number of protein sequences with higher sequence alignment score for each GO term.

The UTMGO and its extended version are available upon request to the corresponding author. The user will be supplied with a set of latest GO RDF/XML files which have been fragmented by the SMAGA, bundled with binary code of the SSMGA and the JAligner engine. Note that, the latest fragmented GO RDF/XML files can be requested depending on the GO monthly releases. Moreover, the user is also advised to have a low-cost PC cluster using MPICH libraries [34] to accelerate the execution speed. Otherwise, the UTMGO and its extended version can be run on a single PC with the following minimum requirements: Fedora Core 2 or Red Hat Linux 8.0 with Pentium 4 processor 2.8 GHz and 512 MB RAM, Linux web browser such as Firefox, Opera, or Konqueror, and Apache Tomcat and JRE (Java Runtime Environment) need to be installed in order to run the JSP page and the JAligner engine respectively.

IV. TESTING AND COMPARISON

GO data released in May 2006 is used to test the UTMGO and its extended version. They are executed using a low-cost PC cluster that consists of 25 Pentium IV 2.8 GHz processors with 512 MB RAM and 100 Mbps NIC. The operating system used is Fedora Core 5. Screenshot of the UTMGO is shown in Fig. 5. The example shows the query results for “DNA binding” (GO:0003677), which is a molecular function term interacting selectively with DNA. The results depict 20 GO terms that have higher term similarity score to the query GO term. The first column presents the GO term accession number, followed by short description of the GO term, its aspect (either cellular component (C), molecular function (F), or biological process (P)), and the term similarity score. Clicking on the respective GO term accession number shows all information related to the term. The information is displayed via AmiGO (www.godatabase.org).

Associated Protein Sequences

Fig. 6 Screenshot of the extended UTMGO with Option 1

Output
The input protein sequence to demonstrate the extended UTMGO is as follows:

```
MVRGKTQMKRIENPTSRQVTFSKRRNGLLKKAFE
LSVLCDAEVALIVFSPRGKLYEFASASTQKTIERYR
TYTKENIGNKTVQDIEQVKADADGLAKKLEALE
TYKRKLLGEKLDECSIEELHSLEVKLERSLISIRGR
KTKLLEEQVAKLREKEMKLRKDNEELREKCKNQP
PLSAPLTVRAEDENPNRTTNDMVDTELFIG
LPGRSRSSGGAEDSQAMPHS
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The protein sequence belongs to AGL20 (MADS box-like protein), an Oryza sativa species that is obtained from Gramene database [35]. The operation of extended UTMGO is divided into two cases: with (Option 1) or without (Option 2) a GO term entered by the user. The first case is depicted in Fig. 6. Same as with the UTMGO, the output consists of the GO term accession number and its short description, aspect, and term similarity score. Value-added information is arithmetic mean (avg), standard deviation (stdev), and the largest value (max) of the sequence alignment score of Ns number of protein sequences that are attached to the GO terms. The user can explore details of the GO term by clicking its accession number. In addition, the Ns number of protein sequences associated with the GO term can be viewed by clicking the arithmetic mean. The protein sequences are displayed with its symbol, data source, evidence, and reference; thence, with sequence alignment score and E-value between them and the query sequence. The second case is depicted in Fig. 7. The GO term proposed by the JAligner engine or selected by the user that is given to the UTMGO is shown at the top of the results table.

To show the capability of the UTMGO, its output is compared with other GO browsers such as AmiGO, GenNav (mor.nlm.nih.gov/perl/gennav.pl), TAIR Keyword Browser (www.arabidopsis.org/), and QuickGO (www.ebi.ac.uk/ego/). Table I tabulates the GO terms returned by those browsers for search of “DNA binding” (GO:0003677). The term similarity score is computed using semantic similarity measure proposed by Razib et al. [32]. The output is arranged based on first 10 search results generated by each browser. The results show that the GO terms with higher term similarity score such as “nucleic acid binding” (GO:0003676), “RNA binding” (GO:0003723), and “ATP binding” (GO:0005524) are associated with protein sequences.
ordered at the top of the list by the UTMGO. The overall performance is shown in Table II for the same input set of 398 molecular function terms, 551 biological process terms, and 93 cellular component terms. Hence, the UTMGO showed a better precision (83.80%) and the QuickGO provided a better recall (74.39%), whereas the GenNav gives the best running time (0.06 seconds).

The comparison between the extended UTMGO and other protein sequence annotation tools are shown in Table III and Table IV. The GOBlet is not in the comparison list due to its service being temporarily unavailable. The results are obtained with AGL20 as the input protein sequence. The arithmetic mean and the largest value of the sequence alignment score for the protein sequences associated with the GO terms are used to analyze those tools. This is due to the fact that quality of the GO terms relies on the similarity between the query protein sequence and the protein sequences associated with them. Thus, higher is better. As depicted in Table 3, almost all GO terms with higher average of sequence alignment score are listed as top 10 GO terms that are returned by the extended UTMGO.

However, GO terms such as “flower development” (GO:0009908) and “cytoplasm” (GO:0005737) are out of the extended UTMGO radar since their term similarity score is 0.9% and 0.6% respectively. These values are lower than the term similarity threshold (1.0%) set for this testing session. Furthermore, as shown in Table 4, all GO terms that have been linked to protein sequence with the highest sequence alignment score (1,153) are returned by the extended UTMGO. Note that, even though GO terms such as “positive regulation of transcription from RNA polymerase II promoter” (GO: 0045944), “DNA bending activity” (GO: 0008301), and “peptidase activity” (GO: 0008233) have maximum sequence alignment score higher than “actin binding” (GO: 0003779) but they are not listed in the top 10 GO terms that are returned by the extended UTMGO. The reason is their average sequence alignment score is lower than the value for “actin binding” (GO: 0003779). To evaluate the performance of the extended UTMGO as compared to other protein sequence annotation tools, a set of protein sequences were chosen from Gramene (www.gramene.org), a database of *Oryza sativa*; Esembl (www.ensembl.org), a database of *Homo sapiens*; SGD (www.yeastgenome.org), a database of *Saccharomyces cerevisiae*; and TAIR (www.arabidopsis.org), a database of *Arabidopsis thaliana*. These protein sequences were selected randomly with 50 protein sequences from each database. The results, as depicted in Table V, show that the extended UTMGO provides a better precision (91.04%) and the GOtcha offered a better recall (89.74%). The best running time is 127 seconds that is taken by the GoFigure.

### Table I

**Comparison of Search Results Between UTMGO with Other GO Browsers**

<table>
<thead>
<tr>
<th>Term</th>
<th>UTMGO</th>
<th>AmiGO</th>
<th>GenNav</th>
<th>TAIR Keyword Browser</th>
<th>QuickGO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Term Accession Number</td>
<td>Term Similarity Score</td>
<td>Term Accession Number</td>
<td>Term Similarity Score</td>
<td>Term Accession Number</td>
<td>Term Similarity Score</td>
</tr>
<tr>
<td>GO:0003677 52.1%</td>
<td>GO:0003677 100.0%</td>
<td>GO:00056092 1.9%</td>
<td>GO:0003681 4.2%</td>
<td>GO:0003677 100.0%</td>
<td>GO:0006260 3.4%</td>
</tr>
<tr>
<td>GO:00053723 24.6%</td>
<td>GO:0003677 100.0%</td>
<td>GO:0019237 4.6%</td>
<td>GO:0003681 4.2%</td>
<td>GO:00051880 2.5%</td>
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</tr>
<tr>
<td>GO:0005524 13.2%</td>
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<td>GO:0031490 3.6%</td>
<td>GO:0003677 100.0%</td>
<td>GO:0003684 11.4%</td>
<td></td>
</tr>
<tr>
<td>GO:0005515 13.0%</td>
<td>GO:0003677 100.0%</td>
<td>GO:0031490 3.6%</td>
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<td>GO:0003684 11.4%</td>
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<td>GO:0003677 100.0%</td>
<td>GO:0003684 11.4%</td>
<td></td>
</tr>
</tbody>
</table>

### Table II

**Comparison of Performance Between UTMGO with Other GO Browsers**

<table>
<thead>
<tr>
<th>UTMGO</th>
<th>AmiGO</th>
<th>GenNav</th>
<th>TAIR Keyword Browser</th>
<th>QuickGO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Precision</td>
<td>83.80%</td>
<td>72.84%</td>
<td>71.62%</td>
<td>72.13%</td>
</tr>
<tr>
<td>Recall</td>
<td>70.35%</td>
<td>67.78%</td>
<td>66.18%</td>
<td>66.96%</td>
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<td>Running time</td>
<td>0.13s</td>
<td>0.09s</td>
<td>0.06s</td>
<td>0.08s</td>
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</tbody>
</table>

V. CONCLUSION

The UTMGO has been presented as an alternative way to search the GO terms. The search is done by finding a group of semantically similar GO terms that relate to the user request. This semantic similarity search is not based on word matching but according to degree of relationships between the GO terms. A gene product that associates with one or more GO terms is used to calculate the amount of information the GO terms share in common. Hence, it gives the degree of relationships. The search results have indicated that the UTMGO is capable to find functionally related GO terms as compared to other existing GO browsers which are based on keyword queries. The applicability of the UTMGO has been shown by its extended version. The extended UTMGO has the ability to annotate anonymous protein sequence. The protein sequences associated with the GO terms that are returned by the extended UTMGO have higher sequence alignment score.
to the query protein sequence. The SSMGA and the SMAGA components in the UTMGO play an important role in accelerating the search. Moreover, the extended UTMGO is not dependent on BLAST and RDBMS, can be used for not-yet-annotated protein sequences, fully based on the GO data, and relate to all categories in the GO.

Future development direction for the UTMGO is applying it to predict protein function and protein-protein interactions. In the case of the extended UTMGO, enhancement includes the ability to support more than one protein sequence per query and to accept DNA sequence as an input.

**REFERENCES**


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