User and Technical manual

for

GOssTo: Gene Ontology Semantic Similarity Tool

Version 1
# Contents

1 Introduction .......................... 3  
   1.1 The idea of GOssTo ................. 3  
   1.2 Contents of the Package .......... 4  
   1.3 This Document .................... 4  
   1.4 Vocabulary ........................ 4  
   1.5 References ........................ 5  

2 Assumptions ......................... 6  
   2.1 Technical Assumptions ............ 6  
   2.2 Input Assumptions ................. 6  
   2.3 Performance ...................... 7  
   2.4 Output considerations ............ 8  
      2.4.1 GOssTo and memory usage ...... 9  
   2.5 Output assumptions ............... 10  

3 Using GOssTo ...................... 11  
   3.1 Using Console Parameters in the standalone version of GOssTo .... 11  
   3.2 GOssToWeb ........................ 12  
   3.3 How to add a new semantic similarity measure ............... 13  
   3.4 Example .......................... 14  
      3.4.1 Step 1. Class definition and constructor .......... 15  
      3.4.2 Step 2. Define the required methods .......... 16  
   3.5 How to add a new extension measure ..................... 17  

4 Walkthrough ....................... 19  
   4.1 Step by Step Guide to Using the Application ............ 19  
   4.2 User FAQ .......................... 24  
   4.3 Using GOssTo as a Library ........ 25  

5 Error Messages .................... 26  
   5.1 Errors Relating to Console Parameters ............... 26  
   5.2 Errors Relating to Input Validation ............... 27  
   5.3 Errors Relating to Calculation and Parsing ........... 28  
   5.4 Other Errors ..................... 29  
      5.4.1 Preamble ...................... 31  
      5.4.2 Terms and conditions .......... 32  

2


1 Introduction

Welcome to the manual for GOssTo; the Gene Ontology Semantic Similarity TOol. The GOssTo application is a practical and fully functional implementation of the ideas set out in Yang et al.’s paper ‘Improving GO semantic similarity measures by exploring the ontology beneath the terms and modelling uncertainty’. The purpose of this software is to compare biological data and produce a meaningful metric describing the likeness or similarity of the biological data in question.

The application will take files from the Gene Ontology Project as input and calculate semantic similarity values based off of six initial, and previously established, ‘host’ similarity measures (HSM). The ‘integrated similarity measure’ or ISM will then use the results of the HSMs and calculate a more meaningful value using the formulae set out in the aforementioned paper.

1.1 The idea of GOssTo

GOssTo was created as a standalone tool to provide a reference implementation for the Random Walk-based measure of Yang et al. Given the fact that Yang’s approach requires a “host” similarity measure (HSM) as the starting point, we also implemented a few of the most popular semantic similarity measures, such as Resnik, Lin, Jiang, etc. This will allow the user two things:

1. To run one of HSMs for a particular annotation of an organism and a certain version of the Gene Ontology.

2. To run Yang’s et al. ISM over the same data.

Our idea, by providing a standalone tool, was to give the user the opportunity to run their own experiments providing his/her GOA files and preferred version of the Gene Ontology, allowing the selection of large variety of different parameters. Given the way ISM was conceived, we were also able to compute HSMs without the need to obtain the improved similarity measure.

As stated, it is clear that GOssTo does not pretend to be an exhaustive implementation of all known semantic similarity measures, as it is conceived as a proof of concept implementation of Yang’s et al. Random Walk Contribution measure.

We have designed GOssTo as an extensible tool. Users can add new similarity semantic similarity measures and add-ons, taking advantage of the large code base available. In order to do that, we have added a section in entitled “How to implement your own semantic similarity measure” which provides detailed instructions on how to extend
GOssTo. If you have developed a new similarity measure for GOssTo, please submit it to support@paccanarolab.org, we will test it and include it in the next release.

1.2 Contents of the Package

The GOssTo package contains an executable .Jar file encompassing all the source files for the program and from which the application will be launched. Also contained within the package are a selection of GOA annotation files and a copy of the gene ontology .OBO file from which similarity values can be readily calculated. These files may need to be updated from the Gene Ontology Project website (http://www.geneontology.org/GO.downloads.shtml) as they are regularly updated with new data. Using out of date files may lead to the generation of inaccurate results.

1.3 This Document

The purpose of this document is to aid the proper use of and provide a comprehensive understanding of the GOssTo application that will lead users to become professional users of the package. This document will offer detailed information on any and all assumptions made in the construction of the program, information regarding licensing, details of planned or developed extensions to the software and a full walkthrough and technical detail section to aid users getting to grips with the system.

1.4 Vocabulary

This section further details some of the more complex technical language or abbreviations used within this document.

- **GOssTo**: The abbreviated name of this software package, standing for: Gene Ontology Semantic Similarity TOol.
- **OBO**: OBO is the name of the flat file format used by the Gene Ontology project to store the central ontological data. The format uses a tag structure with the simultaneous goals of being easily parsed and readable by humans.
- **GOA**: GOA is the name of the file format or plain text structure used for the storage of Annotation files. Unlike OBO, the GOA format is not meant to be read by humans, and so all annotations data is represented by a single line per annotation, with individual data points separated within by tab spaces.
- **HSM**: The abbreviation HSM stands for 'Host Similarity Measure', and refers to previously established formulae for semantic similarity.
- **ISM**: The abbreviation ISM stands for 'Integrated Similarity Measure' in reference to the measure proposed by Yang et al.
• **IO**: An abbreviation for Input Output.

• **Genewise**: A term referring to genes as the base data for calculation.

• **Termwise**: A term referring to GO terms as the base data for calculation.

### 1.5 References

Detailed below are references to any papers or works mentioned or directly referenced within this document and also to the websites of any related or helpful material.

**Papers:**


**Websites:**

- **Gene Ontology Project**: http://geneontology.org/
- **GNU Licenses**: http://www.gnu.org/licenses/
- **Paccanaro Lab**: http://www.paccanarolab.org/
- **Semantic Similarity Paper (Yang et al.)**: http://bioinformatics.oxfordjournals.org/content/28/10/1383
2 Assumptions

2.1 Technical Assumptions

There are several technical assumptions that have been made in relation to the development of this application. As such, all details with respect to the programming language, version numbers and associated packages used within the application are listed below along with the specifications of the development environment. If you have any problems running GOssTo, please first check the data listed below against your own software or hardware and mention in if you need to contact the developers.

Language and Package Details:

- **Programming Language**: Java, version 1.6.0_24
- **IDE**: Eclipse Juno, version 4.2
- **External Packages**: JAMA, version 1.0.3

Development Environment:

- **OS**: Xfce 4.8 64bit
- **RAM**: 3.8GB
- **CPU**: Intel Core i5 CPU (4 x 2.67GHz)
- **CPU Cache**: 8162KB

2.2 Input Assumptions

All input for the application will be taken either by command line parameters upon initiating the application from the Jar file, or requested by the application once it has started. These inputs will take the form of file paths for the OBO and GOA files, text input of GO terms or gene IDs and numerals or booleans representing choices presented by the application.

The inputs that the application offers the user are as follows:

- File path location for OBO and GOA files
- Choice of evidence codes
- Choice of Gene Ontology relations to be used
• Choice of HSM measure
• Decision of whether or not to compute the ISM for the selected HSM
• Which ontology(s) to utilise
• Whether or not to do term based or gene based similarity
• Names for the output files
• Choice of using individual GO terms or Gene IDs

More information regarding the OBO and GOA file formats can be found here: http://geneontology.org/GO.contents.doc.shtml#file

2.3 Performance

The length of computation time naturally varies dependent on the choice of HSM, whether or not an ISM is to be computed, how many files are to be output and the size of the GOA annotation file being used for the calculation.

For the calculation of just an HSM, it should take in the region of 1.5-2 minutes dependent on the choice of annotation file. For an ISM it should take between 6.5-7.5 minutes. For gene based similarity calculations, the time taken is entirely dependent on the number of genes you wish to calculate the semantic similarity for, the less genes the quicker the computation.

When tested using the Yeast annotation file for term-wise semantic similarity calculation, the application took 83 seconds to calculate solely the HSMs and 6 minutes and 7 seconds to calculate the ISMs.

With respect to memory requirements, they are entirely dependent on the number of terms or genes considered in the calculation. Given that we store similarities using single-precision floating point matrices, every cell in the matrix would require 4 bytes.

Memory usage for a few organisms is presented, in GigaBytes, in the following table:

<table>
<thead>
<tr>
<th>Organism</th>
<th>Termwise</th>
<th>Genewise</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arabidopsis</td>
<td>0.24</td>
<td>1.85</td>
</tr>
<tr>
<td>Rat</td>
<td>0.48</td>
<td>0.52</td>
</tr>
<tr>
<td>Mouse</td>
<td>1.14</td>
<td>5.72</td>
</tr>
<tr>
<td>Fly</td>
<td>0.28</td>
<td>0.77</td>
</tr>
<tr>
<td>Yeast</td>
<td>0.26</td>
<td>0.60</td>
</tr>
</tbody>
</table>

To increase the memory allocated use the -Xmx parameter when launching the execution. From Java’s manual (http://docs.oracle.com/javase/6/docs/technotes/tools/windows/java.html):

-Xmxn Specify the maximum size, in bytes, of the memory allocation pool.
This value must a multiple of 1024 greater than 2MB. Append the letter k or
K to indicate kilobytes, or m or M to indicate megabytes. The default value
is chosen at runtime based on system configuration. For more information,
see HotSpot Ergonomics Examples: -Xmx83886080 -Xmx81920k -Xmx80m

The following example is based on the GOssTo and the example data available at:
http://www.paccanarolab.org/gossto/. To run the example with 3 GigaBytes of mem-
ory, we modify the command as follows: ava -Xmx3000 -jar Gossto.jar -calculationdata
termwise -calculationtype ism -evidencecodes EXP,IDA,IP1,IMP,IG1,IEP,TAS,IC -goapath
gene_association.goa_yeast -obopath gene_ontology_ext.obo -hsm Resnik -hsmoutput demo-
hsm_output -ismoutput demo_ism_output -ontology all -relations is_a,part_of -weightedJaccard
true -terms all

This would make 3 GB of memory available to GOssTo. Keep in mind that the
total memory in the system is shared by all running applications, so increasing the
available memory for GOssTo might have an impact on the overall performance of
your system. For larger computations, you might want to consider using GOssToWeb:
www.paccanarolab.org/gosstoweb

These figures are acknowledging that the application calculates the similarity for all
three ontologies in each instance of its running.

2.4 Output considerations

It is important to clarify the nature of the results in light of the semantic similarity
calculations performed by GOssTo. Particularly, given that there are combinations of
parameters that might result in a matrix of zeroes.

To explain this phenomenon, we will resort to the well-known Resnik similarity mea-
sure.

Resnik computes the similarity of two nodes in an ontology by computing the informa-
tion content of the lowest common ancestor of two given nodes. By lowest common
ancestor we mean the first ancestor node shared by both nodes in question.

We define information content of the node as the minus 10 logarithm of the fraction of
genes annotated by that node and the total number of annotated genes, that is:
\[ -\log_{10} \frac{\text{#genes\ in\ node}}{\text{#total\ genes}} \]. The information content will be zero every time the fraction
\( \frac{\text{#genes\ in\ node}}{\text{#total\ genes}} \) is 1. This is equivalent to say that there is, in fact, no infor-
mation content whatsoever in a node that annotates all genes.

The true-path-rule states that every gene annotated by a particular node in the Gene
Ontology is also annotated by all ancestors of that particular node. This means that the
root of the GO DAG will annotate all genes in that particular ontology, which means
that the number of genes annotated by the root of the GO DAG will be the same as the
total number of annotated genes.

Therefore, everytime the user obtains a matrix composed of zeroes, the genes repre-
sented in said matrix did not share any ancestors other than the root of the ontology
itself. We illustrate this case in the following figure:

Figure 2.1 represents a toy example, in which only four nodes of the Gene Ontology
are annotated. Consider two distinct cases, the first one represented by the blue circles
Figure 2.1: Lowest common ancestors and information content

(nodes a and b) and the second one represented by the orange circles (nodes c and d). The colour lines, blue and orange respectively, represent the true path from each node.

If we first analyse the blue case, by following the blue lines we can reach the lowest common ancestor, represented by the node $LCA(a, b)$. In this case, the number of genes annotated by the lower common ancestor of nodes a and b will definitely not annotate all the genes. In fact, the genes annotated by the orange node d are not annotated by $LCA(a, b)$ . This means that the fraction, $\frac{\# genes_{in\_node}}{\# total\_genes} \neq 1$ will definitely be different than 1 for the node $LCA(a, b)$, leading to a non-zero information content and thus to a non-zero Resnik similarity.

The other case, represented by the orange circles, we can, by following the orange lines in this case, reach the lowest common ancestor of nodes c and d. This node, represented by node $LCA(c, d)$, happens to be the root node of this example graph. The fact that this node is the root means that, due to the true path rule, it is in fact annotating all genes. This results in $\frac{\# genes_{in\_node}}{\# total\_genes} = 1$, which means that the information content of it is, effectively, zero.

The user must, therefore, not be surprised in the cases in which he obtains a matrix composed exclusively of zeroes, for it simply means that the nodes do not share relevant information.

2.4.1 GOssTo and memory usage

GOssTo can potentially use a lot of RAM memory. Most of the calculations are done in memory and the result is kept there until printed, for efficiency reasons. Therefore, it is likely that some of the “large” eukaryots (cow, human, etc.) will require a lot of memory, especially while doing genewise computations. We have, however, tried to reduce the memory footprint to a minimum, doing all the computations in single-precision floating point. Nevertheless, in some special cases, an error like this could be shown:
ERROR: GOssTo has run out of memory: the memory allocated is insufficient for this calculation. To increase the memory allocated use the -Xmx parameter when launching the execution. For example use: java -Xmx3000M -jar gossto.jar to increase the maximum available memory to 3 Gb. Alternatively, please use the web service at www.paccanarolab.org/gosstoweb

Please refer to section 2.3 for a detail explanation on how to address this issue.

2.5 Output assumptions

The output produced by the package is in the form of plain text files. Each file is named by the user and contains an N by N matrix of decimal similarity figures where N is the number of GO terms or Genes used in the calculation. The similarity values are separated by commas and there is an axis of terms, GO ids or Gene ids, along the top and down the left hand side. This allows the output to be both humanly readable and easily parsable by other applications. However, for results files involving many terms it readability will suffer somewhat due to the quantity of numbers and the proximity.
3 Using GOssTo

3.1 Using Console Parameters in the standalone version of GOssTo

The command line application offers the ability to enter all parameters the application would normally prompt for at the point of execution. This increases the speed at which the program can be used by removing this prompt section and starting the calculations straight away.

To utilise this feature, in the line with which you execute the application’s .Jar file, after typing the name of the .Jar simply enter the parameters by typing the parameter label in the form of ‘-parameter’ followed by a space and the appropriate parameter value. The available parameters are listed below:

**Administrative Parameters:**
These parameters takes no arguments; you don’t follow them with any values.

- **-help:** The help parameter brings up a list of all the parameters with their descriptions.
- **-version:** Displays the version of the GOssTo release.
- **-getw:** Displays the section of the licence pertaining to the warranty, or lack thereof.
- **-logfile:** Creates a log file containing notes on the various stages of completion of the GOssTo execution. Used in addition to a full set of calculation parameters.

**Calculation Parameters:**
These parameters DO take arguments; you MUST follow them with the appropriate values described below.

- **-obopath:** Follow this label with the file path of the OBO file you wish to use, no spaces are allowed in this file path.
- **-goapath:** Follow this label with the file path of the GOA file you wish to use, no spaces are allowed in this file path.
- **-relations:** Follow this label with the Gene Ontology relations you wish to use in the calculation. If more than one, separate by a comma only. No Spaces.
• **evidencecodes**: Follow this label with the evidence codes you wish to utilise. Again comma separated and no spaces.

• **hsm**: Follow this label with the name of the HSM you wish to use. The name of the HSM will be case sensitive so please enter the name of the HSM exactly as its class name in its java file.

• **ontology**: Follow this label with either 'hp' to use the Biological Process ontology, 'mf' for Molecular function, 'cc' for Cellular Component or 'all' for all three. Do not enter the inverted commas.

• **calculationtype**: Follow this label with simply 'HSM' or 'ISM' depending upon which you would like to calculate.

• **calculationdata**: Follow this label with 'termwise' to calculate semantic similarity by GO terms, or 'genewise' to calculate by gene.

• **terms**: Follow this label with the ID’s of any terms or genes you wish to focus the calculation on, comma separated. If you do not want to use any specific terms, follow the label with 'all', without the inverted commas.

• **hsmoutput**: Follow this label with the name you wish to entitle the HSM results file with. Please do not put a file extension in your name, result files will be '.txt' text files by default.

• **ismoutput**: Follow this label with the name you wish to entitle the ISM results file with. Please do not put a file extension in your name, result files will be '.txt' text files by default. This field is OPTIONAL if calculating solely HSM values.

**Usage Example:**
The following is an example of the correct entering of the console parameters:

```bash
Usage Example:
The following is an example of the correct entering of the console parameters:
```

If you receive any error messages when using the console parameters, they can be referenced to the Error Messages section at the end of the manual. Suggested action is listed for each error.

### 3.2 GOssToWeb

GOssToWeb provides an intuitive and powerful web interface for GOssTo. Available at [http://www.paccanarolab.org/gostoweb](http://www.paccanarolab.org/gostoweb), it allows extensible configuration of the experiment and provides features such as:
• Email notification
• Full parameter validation
• Up to date organism annotation files and ontologies.

GOssToWeb provides an ideal tool for occasional calculations. We keep the version of GOssToWeb consistent to the provided Java version to ensure identical results.

3.3 How to add a new semantic similarity measure

We will use as a guide the implementation of the Resnik measure which can be found in the file Resnik.java of the HSM package.

1. Download the last version of the source code from: https://github.com/pwac092/gossto.

2. Add a new Java class in the package HSM. The name given here will be the name used to identify the measure from the command line tool (so, the "class Resnik" in "Resnik.java" is referred to as "Resnik", if you want to implement a "class ImprovedResnik", you will be able to invoke it using "ImprovedResnik").

3. Inherit from class "HSM", and complete the required methods. The constructor will probably remain empty.

4. Now, we need to implement two methods: the termwise semantic similarity (i.e., the similarity between two terms) and the genewise semantic similarity (given two genes, which is the semantic similarity between them).

5. To implement the termwise semantic similarity, we are only given a parameter, an integer showing the ontology (0=BP, 1=MF, 2=CC). Note that we assume that the three ontologies are independent, and we will ignore inter-ontology relationships. The structure matrixAxis is a double array of GOTerm objects, which is first indexed by ontology, and secondly by an integer in the interval 0 to numGOtermsPerOntology[ontology] (without including this last one). Once retrieved, every GOTerm object can provide access to the GO term identifier, a pointer to the ontology, the parents and the descendants for each one of the relationships, and other miscellaneous data. In order to store the computed semantic similarity values, we will have to create an NxN Matrix, where N is the number of terms of this ontology. Therefore, we will iterate over every pair of GOTerms, using the two nested for loops, and computing the semantic similarity using this measure. We have provided a function lowestCommonAncestor, implemented in the HSM class, which returns the number of annotations given to the lowest common ancestor in the specified ontology and annotation file, divided by the maximum number of annotations of that ontology.

6. Most of the genewise implementations only require, for every two genes, the computation of a function of all the pairwise values of the termwise Gene Ontology of
the terms associated to both genes. Given the fact that this is independent of the proper implementation of the measure, we have implemented a method "geneWiseSimilarityByMaximum" which computes the maximum of all values given by all possible pair of GO terms where the first one is associated to the first gene and the second one to the second. If you prefer to compute the average or other aggregate function, you can implement your own method using the mentioned one as a base. Please note that this method is optimized and makes use of some optimizations, finding which rows dominate others, hence saving computation time. If you use another measure be careful of not using this optimization, as it may not suit your needs.

Once GOssTo has been recompiled, the new HSM will be available for use. To use this new HSM; when prompted by the program for your choice of HSM, input the name of the new HSM and the application will look for it within its classes and if found, will use it as the HSM for the upcoming calculation.

Note that if your HSM does genewise semantic similarity calculations, it will be told to use the specialised gene ISM implementation strategy. If you do not wish to use this, follow the instructions to change the source code above and in the source code itself.

If you have developed a new similarity measure, you can submit the GOssTo-friendly version of the implementation, and we will do our best to evaluate it quickly and include it in the next release of GOssTo.

There is one last consideration. While the code we provide is licenced under GPLv3, licensing of individual component packages have to be respected.

### 3.4 Example

In order to clarify the previous explanation, we will present an example based on Resnik’s Semantic similarity measure.

```java
public class Resnik extends HSM {

    public Resnik(GOTerm[] allTerms, String[] genes, String[][] goIds,
                  GOTerm[][] axis, String[] targets, Assignment annotations,
                  String[] relations, TinyLogger logw) {

        super(allTerms, genes, goIds, axis, targets, annotations,
              relations, logw);
    }

    @Override
    public Matrix calculateTermWiseSemanticSimilarity(int ontology)
    throws IOException, OutOfMemoryError {

        assert (ontology >= 0 && ontology < 3);
    }
}
final int N = numGOtermsPerOntology[ontology];

Matrix result = new Matrix(N, N);

for (int i = 0; i < N; i++) {
    for (int j = i; j < N; j++) {
        float res = (float) -Math.log(
            lowestCommonAncestor(
                matrixAxis[ontology][i].getAncestors(),
                matrixAxis[ontology][j].getAncestors(),
                ontology)
        );
        result.set(i, j, res);
        result.set(j, i, res);
    }
}

return result;

@Override
public Matrix calculateGeneWiseSemanticSimilarity(int ontology)
    throws IOException, OutOfMemoryError {

    return super.geneWiseSimilarityByMaximum(ontology);
}

This code excerpt presents the complete implementation of Resnik's semantic similarity measure. We will explore it step by step, to illustrate the simplicity of GOssTo's extension procedures. There are no specific input assumptions other that the required structures, and the functions you implement have to return the results in matrix form, as shown in the example. You can implement the measures as you see fit, as long as the end product conforms to this standard.

3.4.1 Step 1. Class definition and constructor

Every new measure has to extend the HSM class. That is, each measure will implement their own specific methods and functions to produce the results, while using existing utilities to do so.

public class Resnik extends HSM {

    public Resnik(GOTerm[] allTerms, String[] genes, String[][] goIds,
In this particular case, we had no need to initialise any of the parent’s class properties, but if the measure you design requires you to, you are free to modify them.

### 3.4.2 Step 2. Define the required methods.

Since the parent class does not actually implement any measure but defines nevertheless placeholders for the methods, you need to override the following methods:

```java
public Matrix calculateTermWiseSemanticSimilarity(int ontology)
public Matrix calculateGeneWiseSemanticSimilarity(int ontology)
```

These methods will implement the actual measures. In this particular case (that is for Resnik’s similarity) we need to determine the Lowest common ancestor of every two nodes, and compute it’s information content. Given that several measures require the Lowest Common Ancestor to be computed, we provide a method implemented in the the parent class.

```java
@Override
public Matrix calculateTermWiseSemanticSimilarity(int ontology)
    throws IOException, OutOfMemoryError {
    assert (ontology >= 0 && ontology < 3);
    final int N = numGOtermsPerOntology[ontology];
    Matrix result = new Matrix(N, N);

    for (int i = 0; i < N; i++) {
        for (int j = i; j < N; j++) {
            float res = (float) -Math.log(
                lowestCommonAncestor(
                    matrixAxis[ontology][i].getAncestors(),
                    matrixAxis[ontology][j].getAncestors(),
                    ontology)
            );
            result.set(i, j, res);
            result.set(j, i, res);
        }
    }
    return result;
}
```
The last method that needs to be implemented is the one that computes the genewise similarity. For Resnik it is simply the maximum of all similarities between any two nodes in the Gene Ontology. Again, since several measures follow a similar strategy, we have implemented this method in the parent class.

```java
@Override
public Matrix calculateGeneWiseSemanticSimilarity(int ontology)
throws IOException, OutOfMemoryError {
    return super.geneWiseSimilarityByMaximum(ontology);
}
```

Once the new class is compiled and properly located, it will become available in the HSM menu in the Interactive interface of GOssTo. This is done through the Java Reflection engine, and the path for the compiled classes is fixed to the same place where the existing measures are located.

### 3.5 How to add a new extension measure

We have designed GOssTo as an open system, in which the different components can be disassembled. This allows the user to modify components in order to take advantage of the provided utilities.

As an example, we will show how to replace the Improved Semantic Similarity for another add-on measure that can extend Host Similarity Measures.

The class

```java
public class ISM_validImplementation
```

implements the add-on measure.

This class does not extend any other class in the project, with which we obtained a decoupled module that can easily be replaced.

The class constructor has to be appropriately initialised as follows.

```java
public ISM_validImplementation(GOTerm[] ISM_currentGoTerms,
                                Matrix HSM,
                                String[] ISM_currentRelations,
                                Assignment ISM_Annotations,
                                boolean termwise,
                                boolean wJaccard,
                                TinyLogger logger)
```
This example is based on ISM, so your measure might not require all these parameters. The input considerations for the different parameters is as follows.

- **GOTerm[,] ISM_currentGoTerms**

- **Matrix HSM**: Contains the Host Similarity values, e.g. in case Resnik was selected, this matrix will contain all the Resnik similarity values.

- **String[,] ISM_currentRelations**: A string array containing the relations to consider in the ontology.

- **Assignment ISM_Annotations**: An object that contains a mapping between GOterm and the annotated genes. The class Assignment defines the object.

- **boolean termwise**: Boolean value that determines whether we compute genewise or termwise similarities.

- **boolean wJaccard**: Boolean value that selects a specific weighting we require for ISM.

- **TinyLogger logger**: A logger object. This is clearly defined in the TinyLogger class.

While we cannot determine whether you will require more values for the function, the only requirement is that the calling function should be extended to include all the parameters.

All output values are expected to be stored in a Matrix object.

In our case, we have implemented the function

```java
private Matrix getISM()
```

which returns the end result of the add-on measure. These methods could be replaced or fully renamed to incorporate the new measure.
4 Walkthrough

4.1 Step by Step Guide to Using the Application

This section aims to give a full walkthrough for an average use of the system. This will be shown using the command line application.

1. The first step is to execute the program itself, this is done by navigating on the console to the location of the GOssTo .Jar file and executing it by calling `java -jar GOssTo.jar` as shown below:

   ![Image of terminal output]

2. Once entered the application will start and prompt you for the location a Gene Ontology Project .OBO file and also that of the Gene Ontology Project GOA file. You will need to enter the full file path for both as shown below:

   ![Image of terminal output]

3. Now that we have entered the source files we are working with, the application will begin to prompt us for the parameters of the calculation. Firstly we are given the option to choose whether we wish to enter the Gene Ontology relationships we wish to use or whether to use is_a as default. For the purposes
of this walkthrough we will choose the default option:

4. We are now presented with the option of choosing the evidence codes that will determine which aspects of the annotation file will be read into the calculation process. For this execution we shall use the default selection that is, for note; EXP, IDA, IPI, IMP, IGI, IEP, TAS and IC. This choice is shown below:

5. Before we choose our HSM, we are now asked whether we wish to calculated semantic similarity based upon GO term by GO term calculations or gene be gene calculations; termwise or genewise. For this walkthrough we shall choose termwise:
6. Now that the base parameters have been decided upon, we are given the choice of similarity measure. First we are to decide if we want to select one from a list or enter the name of one we wish to use. Note that if you wish to use your own HSM, you must enter it by name. For this walkthrough we will select one from the list:

7. The next choice to make is the ontology we want to work with:
8. The application will now want to know if we wish to work with particular terms or whether we wish to calculate the similarity between all terms available. For this walkthrough let us calculate them all:

9. The following prompt asks us if we wish to calculate the HSM similarity matrices or if we also wish to calculate the ISM similarity matrices. It is important to note however that the HSM similarity matrices that are produced when calculating the ISM will not be the same as the regular HSM matrices, due to a few calculation differences that aid the calculation of the ISM results. We wish, naturally, to calculate the ISM:
10. As we approach the end of the user prompt section, we are lastly asked to enter the names we wish to use for the HSM and ISM matrix files. Note that you may wish to precede the name by the location you wish the file to be created in. Please note also that the files will be output as .txt by default so do not enter a file extension in your name. In this example however, we shall create the files in our current location and therefore only specify the name of the files:

11. Now that the parameters have been entered the calculation will begin and the results will be output to either the location specified, or if not; that of the .Jar file.
No more interaction will be required on the part of the user.

If you have any questions or are uncertain of the running of the application, please refer to the FAQ in the next section.

4.2 User FAQ

This section aims to satisfy any simple questions or concerns you may have with the GOssTo package. Below is a listing of questions and answers pertaining to the application:

Q : The calculation is taking a very long time, should I be concerned?
A : No. Calculations involving many terms or genes naturally involve calculating very large matrices of values. This takes quite some time.

Q : Help! I’ve got an error, what should I do?
A : Please take note of the error message and refer to it in the Error Messages section at the end of the manual.

Q : I want to use an OBO/GOA file from a different provider to the Gene Ontology Project, is this okay?
A : If it is formatted to the respective GO file format then theoretically there should not be a problem, give it a try. However GOssTo was built to use files from the GO Project and so we cannot ensure the proper functioning of files from other sources.

Q : My results file contains infinite values, why is this so?
A : Due to the calculation formulae of certain semantic similarity methods, the similarity between two very dissimilar GO terms or genes will sometimes give an infinite value. In regard to similarity, treat it as a zero.

Q : Why are the HSM matrices produced in a ISM calculation not correct as results for that HSM?
A : When undertaking an ISM calculation, the annotation values are altered slightly so as to accommodate for and speed up operations when calculating the ISM matrices later on. These changes naturally alter the the values of the HSM matrix.

Q : Why is my HSM method not working with the application?
A : This could be for a number of reasons. Firstly check that it implements the HSM package, secondly check that the parameters of the HSM constructor are in order, thirdly check if the java file is in the correct location. If none of these are wrong, please refer any error messages you may have to the Error Messages section.
Q: I don’t like the way you’ve done / I think you’re code here would be better if you did .

A: If you have a suggestion or a criticism please fell free to get in touch. Please also have a read of the GNU GPL and feel very much free to alter the source code in any way you see fit.

Q: I like GOssTo and would like to use it in my program, is there an easy way I could do this?

A: Please refer to the section ’Using GOssTo as a Library’ in the Technical Information section. Please also have a read through the GNU GPL if you are thinking of releasing your program.

Q: Where can I find more info about the developers of this package?

A: http://www.paccanarolab.org/

If your question does not appear above, feel free to ask the developers at: .

4.3 Using GOssTo as a Library

This package can be altered for integration as a library for an external software package. This can be done very simply by the removal of the ISM.java file that contains the software’s main method. Once removed, the methods and classes are free to be interacted with at will, however the following paragraph will detail the advised order of function and classes.

There are three interface classes packaged within the software that are used to access all key functionality of the system. These classes are ’GOTree_Interfacer’, HSM_Interfacer and ISM_Interfacer. These three contain all the desirable functionality with respect to their area of specialty; GOTree deals with all aspects of the source data, HSM deals with retrieving and executing the required similarity measure and ISM focuses on accessing the appropriate ISM class and calculation. It is naturally advised that these classes are accessed in the order that they have been listed here. Note that the output method, should it be required, resides within the ’ioValidation’ class.
5 Error Messages

This section details the error messages that you may encounter when using GOssTo and any subsequent action that should be taken.

5.1 Errors Relating to Console Parameters

ERROR: OBO Path Not Specified: The OBO file path has not been specified when using the console parameters. Ensure you are using the '-obopath' parameter.

ERROR: GOA Path Not Specified: The GOA file path has not been specified when using the console parameters. Ensure you are using the '-goapath' parameter.

ERROR: GO Relations Not Specified: The GO relations have not been specified when using the console parameters. Ensure you are using the '-relations' parameter.

ERROR: Evidence Codes Not Specified: The evidence codes have not been specified when using the console parameters. Ensure you are using the '-evidencecodes' parameter.

ERROR: HSM Not Specified: The HSM measure has not been specified when using the console parameters. Ensure you are using the '-hsm' parameter.

ERROR: Ontology Choice Not Specified: The ontology to be used has not been specified when using the console parameters. Ensure you are using the '-ontology' parameter.

ERROR: It Has Not Been Specified Whether an HSM or ISM is Being Calculated: Whether we are calculating an HSM or an ISM has not been specified when using the console parameters. Ensure you are using the '-calculationtype' parameter.

ERROR: It Has Not Been Specified Whether Gene or GO Terms are to be Used: Whether we are doing a termwise or genewise calculation has not been specified when using the console parameters. Ensure you are using the '-calculationdata' parameter.

ERROR: Terms (Gene IDs or GO Terms) Not Specified or Not Set To “all”: The terms (genes or GO terms) to be used for the calculation have not been specified when using the console parameters. Ensure you are using the '-terms' parameter, if you wish not to use any specific terms, add 'all' without inverted commas after -terms.
ERROR: HSM Output Path Not Specified: The HSM output name/location has not been specified when using the console parameters. Ensure you are using the '-hsmoutput' parameter.

ERROR: ISM Output Path Not Specified: The ISM output name/location has not been specified when using the console parameters. Ensure you are using the '-ismoutput' parameter.

ERROR: HSM/ISM specification has been entered incorrectly: The calculation type was not either 'hsm' or 'ism'. Ensure that the '-calculationtype' parameter is followed only by 'ism' or 'hsm' in either upper or lower case.

ERROR: Genewise/Termwise specification has been entered incorrectly: The calculation data option was not either 'termwise' or 'genewise'. Ensure that the '-calculationdata' parameter is followed only by 'termwise' or 'genewise' in either upper or lower case.

ERROR: Parse failed: A problem has occurred relating either to the execution of the program or with the parameters you have used, please check that you are executing the program correctly and that all parameters are present and correct.

5.2 Errors Relating to Input Validation

ERROR: Invalid Entry: The entry you have just made is either: not an available option or the wrong type of data.

ERROR: Invalid Choice: The choice you have made was not one that is available.

ERROR: Inputted relation does not exist; try again: The GO relation you have entered is not a valid GO relation, please enter a valid GO relation.

ERROR: Inputted relation has already been input: The GO relation you have entered is one that has already been entered. Do not enter the same relation more than once.

ERROR: Choice of Ontology Invalid: The choice of ontology you have entered is not a choice that is available, please check your entry an try again.

ERROR: Relation: *specific relation*, is not acceptable or does not exist: The GO relation you have entered is not a valid GO relation, please check your entry and try again.

ERROR: Evidence code: *specific evidence code*, is not acceptable or does not exist: The evidence code you have entered is not a valid evidence code, please check your entry and try again.
5.3 Errors Relating to Calculation and Parsing

ERROR: GOssTo has run out of memory: the memory allocated is insufficient for this calculation. To increase the memory allocated use the -Xmx parameter when launching the execution. For example use: java -Xmx3000M -jar gossto.jar to increase the maximum available memory to 3 Gb. Alternatively, please use the web service at www.paccanarolab.org/gosstoweb.

ERROR: Too many indirection levels found for GO term id: *specific GO ID*: This means that this term is obsolete and points to another term which is also obsolete and either no alternative terms have been processed for it, or, there is more than a level of indirection. Update your OBO file, comment this entry out by placing exclamation marks at the start of each row containing its data or remove this entry.

ERROR: consider tag present for: *specific GO ID*: This error raises attention to the fact that 'consider' tags have been used in the definition of this term. This will not stop calculations taking place but may affect them. To find out more about consider tags refer to the optional tags section on this page: http://geneontology.org/GO.format.obo-1.2.shtml.

ERROR: The kind of relations that are going to be parsed should be set first via GOTerm .setRelations: A GOTerm instance has been created without a relation parameter, check the integrity of your OBO file.

ERROR: The relation *specific relation* is not being parsed in this instance of the Gene Ontology: The relation in question is unrecognised or not accepted, please check your OBO file.

ERROR: GO term doesn’t exist: A GO term is being used that has not been recognised. Please check your OBO file is correct and up to date.
ERROR: Gene IDs entered could not be found in the GOA file supplied: One or more gene IDs is being used that is not recognised. Please check your GOA file is correct and up to date.

ERROR: Problem with source data, please check OBO and GOA files are correct: A problem has arisen when attempting to manipulate the source data. Check that your GOA and OBO files are correct and try again.

ERROR: class *class name* is not instantiable: There was an error in accessing the HSM class in question. Please check that it is in the correct location, that it implements the ‘HSM’ interface and that it is defined and formatted correctly.

ERROR: class *class name* has an inaccessible constructor: The constructor for the HSM class in question is inaccessible by GOssTo, please ensure that its permissions are set to public.

ERROR: class *class name* not found: The HSM class in question could not be found. Please ensure that you typed the name correctly and that the file for that class is in the correct location.

ERROR: class *class name* could not be invoked: There was a problem invoking the HSM class in question. Please ensure that the code for this class is correct and parsable.

ERROR: class *class name* has an illegal argument: This error was triggered by an illegal argument the HSM class in question. Please ensure that the code for this class is correct and parsable.

ERROR: class *class name* has triggered a security exception.: The HSM class in question has caused a security violation. Please ensure that the code for this class is correct and parsable.

ERROR: Problem with the Gene Ontology file.: The Gene Ontology file is possibly corrupted. Term GO:000xxxx points to term GO:000yyyy using the ‘is_a’ relationship. However, the term GO:000yyyy does not appear in your Gene Ontology file (probably it is obsolete). If the error persists with a newer Gene Ontology file, substitute the reference to GO:000yyyy in the term GO:000xxxx y one of the alternatives provided in its ‘obsolete term’ entry.

ERROR: An unexpected error has occurred. Error Signature: *error message or exception listing*: An error has occurred that has not been anticipated, the cause is uncertain. Please refer to the ‘Other Errors’ Section.

5.4 Other Errors

In the unlikely event that an error is triggered that is not listed here, please contact the developers with a screenshot of the error message and instructions on what you have
done to produce this error.
If it is possible to recreate the error, then it would also be very useful if you could
recreate it using the '-logfile' console parameter and supply the resultant file 'LOG.txt'
with your message. All help in solving any bugs will be much appreciated.
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