# API Documentation

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# August 26, 2018

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VariablesModule enrich

#### Module enrich 1

GSEA script. Launches Enrich.R script and manage its results.

### Requires:

 $\bullet$  python 2.7<sup>1</sup> or greater

• R 3.3.3 RC<sup>2</sup>

 $\bullet$  Conda  $4.4.10^3$  or greater with FastGSEA conda environment (fastgsea.yml)

Version: 3.0

Author: Kevin Durimel

License: GPL

#### **Functions** 1.1

launchGSEA(outputDir)

## 1.2 Variables

Name	Description	
doc	Value:	
credits	Value: ['Kevin La', 'Arnaud Felten', 'Meryl	
	Vila-Nova', 'Nicolas	
maintainer	Value: 'Kevin Durimel'	
email	Value: 'k@durimel@gmail.com'	
status	Value: 'Alpha'	
EXEC_DIR	Value:	
	'/home/wrecker/EspaceDeTravailLinux/DNAlogy/aug1	8/DNAlogy
package	Value: None	

 $<sup>^{1} \</sup>rm https://www.python.org/downloads/$ 

<sup>&</sup>lt;sup>2</sup>https://www.r-project.org/ <sup>3</sup>https://conda.io/

Variables Module fastGSEA

# 2 Module fastGSEA

FastGSEA workfow. Performs GO-terms enrichment analysis between two gene sets. These gene sets must be provided as files containing one international databank (ncbi, refseq, etc...) gene or protein identifier per line.

### Requires:

• python  $2.7^4$  or greater

 $\bullet$  R 3.3.3 RC<sup>5</sup>

• Conda 4.4.10<sup>6</sup> or greater with FastGSEA conda environment (fastgsea.yml)

• map.py

• managefiles.sh

Version: 3.0

Author: Kevin Durimel

License: GPL

### 2.1 Functions

$\mathbf{get}\_\mathbf{parser}()$
Initlializing arguments.
Return Value arguments values
(type = object)

 $\mathbf{main}()$ 

### 2.2 Variables

Name	Description
doc	Value:
credits	Value: ['Kevin La', 'Arnaud Felten', 'Meryl
	Vila-Nova', 'Nicolas
maintainer	Value: 'Kevin Durimel'
email	Value: 'k@durimel@gmail.com'
status	Value: 'Alpha'
package	Value: None

 $<sup>^4 \</sup>rm https://www.python.org/downloads/$ 

<sup>&</sup>lt;sup>5</sup>https://www.r-project.org/

<sup>&</sup>lt;sup>6</sup>https://conda.io/

Class Timer Module fastGSEA

~ ~	$\sim$ 1	
2.3		l'imon
$\omega$	Class	$\mathbf{Timer}$

object	
	fastGSEA.Timer

Initlialize and manage timers.

### 2.3.1 Methods

init(self)	
xinit() initializes x; see help(type(x)) for signatu	re
Overrides: objectinit extit(inherited documentation	n)

# Inherited from object

$\_\delattr\_\_$	_(),	$\_{ m format}$	(), _	g	etattribu	ıte	$(), \underline{\hspace{1cm}}$ has	h(),	new_	():
reduce	_(),	$\_{\rm reduce}\_$	_ex	$(), _{-}$	repr	_(), _	$\_\_$ setattr $\_$	(),	_sizeof	_(),
str(),	su	bclassho	ok(	)						

# 2.3.2 Properties

Name	Description
lifetime	
Inherited from object	
class	

# 3 Module map

Id mapping script for a gene set. Takes as input a file (gene set) constitued by Refseq, Supported ids are (by decreasing quantity of available information):

- UniProtKB-AC (1st col in gz file)
- 2. UniProtKB-ID (2)
- 3. GeneID (EntrezGene) (3)
- 4. RefSeq (4)
- 5. GI (5)
- 6. GO (as output only) (7)
- 8. UniRef100 (8)
- 9. UniRef90 (9)
- 10. UniRef50 (10)
- 11. UniParc (11)
- 15. UniGene (15)
- 17. EMBL (17)
- 18. EMBL-CDS (18)
- 19. Ensembl (19)
- 20. Ensembl\_TRS (20)
- 21. Ensembl\_PRO (21)

@requires: U{python 2.7<https://www.python.org/downloads/>} or greater

Orequires: U{Conda 4.4.10<https://conda.io/>} or greater with FastGSEA conda environment

Orequires: manageFiles.sh

Version: 3.0

**Author:** Kevin Durimel

License: GPL

#### 3.1 Functions

 $show\_progression(counter, total, precision)$ 

mk\_susbet(idMappingFile)

Functions Module map

### ids\_to\_go(idMappingFile, idsFile, outputPrefix)

Takes as input Uniprot or Refseq ids and returns its corresponding GO-terms. Please use this function if you want to retrieve GO-terms using a input file containing Refseq or Uniprot ids. The id mapping file subset is automatically generated if it was not already the case (e.g First script launch , or file deleted.)

### **Parameters**

idMappingFile: path to the .gz file used for id mapping

(type=string)

idsFile: path to the input file containing all the ids we have

to map

outputPrefix: file path output prefix

(type=string)

view: (type=string)

# ids\_to\_go\_online(idMappingFile, idsFile, outputPrefix)

This function follows same principles as ids\_to\_go() but performs id mapping online. It the best choice for reliability and up-to-date GO-terms.

Ids\_to\_to\_online() always requires idMappingFile parameter to retrieve refseq ids offline. Takes as input Uniprot or Refseq ids and returns its corresponding GO-terms. Please use this function if you want to retrieve GO-terms using a input file containing Refseq or Uniprot ids.

### **Parameters**

idMappingFile: path to the .gz file used for id mapping

(type=string)

idsFile: path to the input file containing all the ids we have

to map

outputPrefix: file path output prefix

(type=string)

view: (type=string)

Functions Module map

# any\_ids\_to\_go(idMappingFile, idsFile, outputPrefix, whichDb)

Takes as input any suppported ids and returns its corresponding GO-terms. Slower than ids\_go\_to beacause it do not use the idMappingFile subset and requests id per id.

### **Parameters**

idMappingFile: path to the .gz file used for id mapping

(type=string)

idsFile: path to the input file containing all the ids we have

to map

whichDb: the ids database prefix that we want in output (for

example, if we want go ids, which DB = GO. Please

refer to "supported ids" at the top section)

(type=string)

outputPrefix: file path output prefix

(type=string)

view: (type=string)

# any\_ids\_to\_go\_online(idMappingFile, idsFile, outputPrefix, whichDb)

This function follows same principles as any\_ids\_to\_go() but performs id mapping online. This function is an experimental feature and support only the ids supported by the Uniprot API<sup>a</sup> Takes as input any supported ids and returns its corresponding GO-terms. Slower than any\_ids\_go\_to() beacause it do not use the idMappingFile subset and requests id per id.

#### **Parameters**

idMappingFile: path to the .gz file used for id mapping

(type=string)

idsFile: path to the input file containing all the ids we have

to map

whichDb: the ids database prefix that we want in output (for

example, if we want go ids, which DB = GO. Please

refer to "supported ids" at the top section)

(type=string)

outputPrefix: file path output prefix

(type=string)

view: (type=string)

 $<sup>^</sup>a$ https://www.ebi.ac.uk/QuickGO/api/index.html

Variables Module map

# 3.2 Variables

Name	Description	
doc	Value:	
credits	Value: ['Kevin La', 'Arnaud Felten',	
	'Meryl Vila-Nova', 'Nicolas	
maintainer	Value: 'Kevin Durimel'	
email	Value: 'k@durimel@gmail.com'	
status	Value: 'Alpha'	
EXEC_PATH	Value:	
	'/home/wrecker/EspaceDeTravailLinux/DNAlo	gy/aug18/DNAlogy.
package	Value: None	

## 4 Module trim

GOtrimmer script. Parse object data from a .obo file, and provide a subset of non-prokaryotic and non obsolete GO-terms based on the lightest official Uniprot's gene ontology file (GO-basic.obo). The file generated by this script will serve for non-prokaryotic GO-terms trimming. Obo files are daily updated and can be found here: http://viewvc.geneontology.org/viewvc/GO-SVN/ontology-releases/YYYY-MM-DD/subsets/ (replace YYYY-MM-DD by currrent date) Please notice that 'gosubset\_prok' terms are not maitained since 2018/06 because they may be irrelevant. More information here: https://github.com/geneontology/go-ontology/pull/16255

### Requires:

• enrich.R output file (hyperesults.csv)

• old GO-basic.obo file (< 2018/06 releases) or gosubset\_prok.obo file.

Version: 3.0

Author: Kevin Durimel

License: GPL

#### 4.1 Functions

$add\_object(d, outputFile)$
Add GO-terms objects only if they are non-prokaryotic and up to date (i.e non
obsolete)

```
mk_subset(oboFile, outputFile)
Generates a GO-terms file subset based on a obo file.
```

 $\mathbf{trim}(goprok\_txt,\ enrichmentResultsFile)$ 

#### 4.2 Variables

Name	Description
doc	Value:
credits	Value: ['PM 2Ring :
	http://stackoverflow.com/q/32989776/4014959
maintainer	Value: 'Kevin Durimel'
email	Value: 'k@durimel@gmail.com'
status	Value: 'Alpha'
all_objects	Value: {}

continued on next page

Variables Module trim

Name	Description
package	Value: None

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