Date: 2010-12-21

# JCVI Metagenomics Annotation Pipeline Process Documentation

Data Snapshot Directory Location: /usr/local/projects/DB/MGX/prok-annotation-pipeline/current

File	Date	Description
ALL_LIB.HMM	12/09/2010	HMMER3 model library collection (contains TIGRFAM and PFAM models)
hmm3.db	12/10/2010	Sqlite database with HMMER3 model meta-information stored in two tables (hmm3 and
		hmm_go_link). Information stored includes Iso-Type, EC, GO, cut offs, etc <b>Used in steps 2.2</b> and <b>2.3</b> .
synonyms.tab	12/21/2010	Tab-delimited file with synonym pairs. Column 1 non-preferred name; Column 2 preferred name.
		The file is used to replace non-preferred with preferred names in the final annotation process.
		Names have been exported from the Protein Naming Utility
		http://www.jcvi.org/pn-utility/web/download/download full matches.php
		Used in step 6.3.
Uniref100.db	11/30/2010	Sqlite database with selected UniRef100 information stored in the annotation table. Column 1
		accession; Column 2 key; Column 3 value. Keys that are stored are gene symbol (GN), GO, EC,
		CAZY, reviewed state (Swissprot/TrEMBL). Used in step 3.3.
Uniref100.tab	11/30/2010	Tab delimited file with comprehensive UniRef 100 cross-reference information. Column 1:
		accession; Column 2: key; Column 3: value. A subset of keys described in
		http://www.uniprot.org/docs/dbxref are available
Uniref100.fasta	11/30/2010	Used as subject database for the blastp searches. Used in step 3.1
		This file contains all UniRef100 entries in FASTA format. The definition line in the FASTA format
		includes cluster specific information such as cluster name, number of members and and common
		taxonomy and also the ID of the representative protein. The format is as follows:
		>UniqueIdentifier ClusterName n=Members Tax=Taxon RepID=RepresentativeMember
		where: UniqueIdentifier is the primary accession number of the UniRef cluster. ClusterName is the
		name of the UniRef cluster. Members is the number of UniRef cluster members. Taxon is the
		scientific name of the lowest common taxon shared by all UniRef cluster members.
		RepresentativeMember is the entry name of the representative member of the UniRef cluster. For example:
		>UniRef100_P99999 Cytochrome c n=5 Tax=Hominidae RepID=CYC_HUMAN

# **Step 1 Split Sequences**

Split sequences for parallel searching (Steps 2-6)

executable	split_multifasta.pl
input	fasta file
output	multiple split fasta files
command	<pre>split_multifasta.plinput_file=input.fastaoutput_dir=/tmp output_list=/tmp/split.listoutput_file_prefix='split_' seqs_per_file=50000compress_output=0</pre>

Date: 2010-12-21

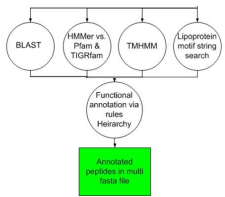


Figure 1 Overview (Steps 2-5 are executed in parallel)

# Step 2 HMMER3 Component (Pfam & TIGRfam)

### Step2.1 Run HMMER3 search

executable	hmmscan
version	HMMER 3.0
data dependencies	formatted ALL_LIB.HMM
input	split fasta file
output	HMMER3 raw output (hmmer3.outr_0)
command	hmmscan -Z 15908 -o <tmp-dir>/hmmer3.outr_0tblout <tmp-dir>/hmmer3.SeqHits.tblr_0domtblout</tmp-dir></tmp-dir>
	<tmp-dir>/hmmer3. DomainHits.tblr_0 <hmm3-db-dir> <input-file></input-file></hmm3-db-dir></tmp-dir>

### Step 2.2 Parse HMMER3 results; generate tab delimited file (JCVI HTAB)

Parses output files generated by hmmscan. Uses sqlite database (hmm3.db) to fetch HMM meta-information (HMM Iso-Type, cut offs, etc.).

executable	htab.pl
data dependencies	sqlite database hmm3.db
input	split HMMER3 raw files (hmmer3.outr_0)
output	split HTAB files (hmmer3.htab)
command	cat hmmer3.outr_0  perl htab.pl -d <snapshot-dir>/hmm3.db &gt; hmm3.htab</snapshot-dir>

### Step 2.3 Parse JCVI HTAB

Performs HMM annotation lookups for common name, gene symbol, GO, and EC assignments from a sqlite database (hmm3.db). Classifies HMM hits based on HMM Iso-Types (10 classes, see box below).

executable	camera_parse_annotation_results_to_text_table.pl
data dependencies	sqlite database hmm3.db
input	JCVI HTAB (hmm3.htab)
output	JCVI HTAB parsed(hmm3.htab.parsed)
command	perl camera_parse_annotation_results_to_text_table.plinput_file hmm3.htabinput_type HTAB
	output_file hmm3.htab.parsedwork_dir <snapshot-dir></snapshot-dir>

```
HMM ISO-TYPES
if ($iso_type =~ /^(equivalog)$|^(PFAM equivalog)$/) {
           $type .= 'Equivalog';
      } elsif ($iso type =~ /^(hypoth equivalog)$/) {
           $type .= 'HypotheticalEquivalog';
      } elsif ($iso_type =~ /^(exception)$/) {
           $type .= 'Exception';
     stype:- makeption;
} elsif ($iso_type =~ /^(subfamily)$/) {
   $type .= 'Subfamily';
} elsif ($iso_type =~ /^(superfamily)$/) {
   $type .= 'Superfamily';
      } elsif ($iso_type =~ /^(equivalog_domain)$|^(PFAM_equivalog_domain)$/) {
           $type .= 'EquivalogDomain';
      } elsif ($iso_type =~ /^(hypoth_equivalog_domain)$/) {
           $type .= 'HypotheticalEquivalogDomain';
      } elsif ($iso_type =~ /^(subfamily_domain)$/) {
   $type .= 'SubfamilyDomain';
      } elsif ($iso type = ~ /^(domain) $/) {
           $type .= 'Domain';
        elsif ($iso_type =~ /^(PFAM)$/) {
  $type .= 'Uncategorized';
        else {
           $type = '';
```

### Step 3 BLAST Component

#### Step 3.1 Run BlastP

Run blastp on individual fasta split files and generate JCVI BTAB format from blast XML output (-m 7 option)

executable	blastall
input	split fast file
output	Blast results in XML format
command	blastall -v 10 -b 10 -X 15 -e 1e-5 -M BLOSUM62 -J F -K 10 -f 11 -Z 25.0 -W 3 -U F -I F -E -1 -y 7.0 -G -1 -A
	40 -Y 0.0 -F "T" -g T -p blastp -z 1702432768 -m 7'

#### Step 3.2 Convert XML files to JCVI tab delimited blast result files (BTAB)

executable	blast_xml_to_btab.pl
input	XML formatted blastp results

output	Tab-delimited blastp results (BTAB)
command	blast_xml_to_btab.pl < blastp.xml > blastp.btab

### Step 3.3 Parse JCVI BTAB

Perform UniRef100 defline lookups (sqlite database uniref.db) for gene symbol, GO, EC, CAZY, and reviewed status (Swissprot or TrEMBL entry). Classifies blastp hits based on sequence coverage and identity (5 classes, see box below).

executable	camera_parse_annotation_results_to_text_table.pl
data dependencies	sqlite database uniref.db
input	JCVI BTAB (blastp.btab)
output	JCVI BTAB parsed
command	perl camera_parse_annotation_results_to_text_table.plinput_file blastp.btabinput_type BTAB output_file blastp.btab.parsedwork_dir <snapshot-dir></snapshot-dir>

# Step 4 Lipoprotein Motif Search

#### Step 4.1 Run lipoprotein motif search

Scans for membrane lipoprotein lipid attachment sites on amino acid sequence. Uses PROSITE motif (^.{0,6}[KR]).{0,18}[^DERK][^DERK][^DERK][^DERK][^DERK][^DERK][LIVMFWSTAG][LI

executable	lipoprotein_motif.pl
input	split fast file
output	BSML formatted file
command	lipoprotein_motif.plinput split1.fastaoutput lipoprotein_out.bsmlgzip_output 0id_repository
	workflow/project_id_repositoryis_mycoplasm 0

# Step 4.2 Parse lipoprotein motif results

executable	camera_parse_annotation_results_to_text_table.pl
input	BSML formatted file (lipoprotein_out.bsml)
output	BSML parsed file (lipoprotein_out.bsml.parsed)
command	camera_parse_annotation_results_to_text_table.plinput_file lipoprotein_out.bsmlinput_type
	LipoproteinMotifBSMLoutput_file lipoprotein_out.bsml.parsed
	/peptide.fasta.q1_q10_1532122841942589727.bsml.parsedwork_dir /tmp

# Step 5 TMHMM Search

Scans proteins for trans-membrane domains.

### Step 5.1 Run TMHMM

executable	tmhmm
version	2.0
input	split fast file
output	tmhmm_out.raw
command	tmhmm split1.fasta > tmhmm_out.raw

### **Step 5.2 Parse TMHMM results**

executable	tmhmm2bsml.pl
version	2.0
input	TMHMM raw file (tmhmm_out.raw)
output	BSML formatted file (tmhmm_out.bsml)
command	tmhmm2bsml.plinput tmhmm_out.rawoutput tmhmm_out.bsmlfasta_input split1.fasta
	compress_bsml_output 0id_repository workflow/project_id_repository

#### Step 5.3 Parse TMHMM BSML

executable	camera_parse_annotation_results_to_text_table.pl
version	2.0
input	TMHMM BSML file (tmhmm.bsml)
output	parsed BSML formatted file (tmhmm_out.bsml.parsed)
command	camera_parse_annotation_results_to_text_table.plinput_file tmhmm_out.bsmlinput_type TMHMMBSML
	output_file tmhmm_out.bsml.parsedwork_dir /tmp

# Step 6 Annotation Rules

The final annotation for each peptide is being derived based on all previously collected evidences. How evidences are being used to assign the various annotation data types (common name, gene symbol, EC, GO, Tigr Role) is based on a evidence rules hierarchy in lib/CAMERA/AnnotationRules/PredictedProtein.pm.

Date: 2010-12-21

# Step 6.1 Concatenate parsed results obtained in steps 2-5

executable	cat
input	all parsed files
output	out.cat.sorted
command	cat *.sorted > out.cat

# Step 6.2 Sort the concatenated file

executable	sort
input	concatenated results (out.cat)
output	sorted concatenated results (out.cat.sorted)
command	sortkey=1,1 -T /tmp -S 1G -d -o out.cat.sorted out.cat

# Step 6.3 Generate tab delimited annotation file (final output)

executable	camera_annotate_from_sorted_table.pl
data dependencies	tab-delimited synonyms.tab file
input	sorted concatenated files (out.cat.sorted)
output	tab delimited annotation results (annotation.tab)
command	perl camera_annotate_from_sorted_table.plinput out.cat.sortedsynonyms <snapshot-dir>/synonyms.tab output out.cat.tmp &gt; annotation.tab</snapshot-dir>