Sequence Similarity Search

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1 BLAST

Homologs are sequences that are related by descent. **Orthologs** are homologs present in two separate species. This does not assume that two homologs are similar in sequence. However, it's a common practice to find orthologs using sequence similarity.

One of the simple ways to find orthologs of a protein from one species in another is to find the protein's **best hit** in the second species and consider the hit as ortholog only when the the hit itself also has the starting protein as the best hit. This relationship is called "Reciprocal Best BLAST hit" or RBH.

We will use RBH to determine all the orthologs between *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe*. This requires all proteins in this two genomes to be searched against all other proteins. This type of BLAST searches are commonly called **all vs all** BLAST.

1.1 Downloading and preparing the input files

First let's create the SC proteome file.

```
# Create a directory for SC
mkdir -p SC

# Change to that directory
cd SC
```

```
# Download all the files of SC from NCBI
wget --quiet ftp://ftp.ncbi.nih.gov/genomes/Fungi/Saccharomyces_cerevisiae_uid128/*.faa

# Combine all the .faa files into one file
cat *.faa >sc.fas

# We no longer need the .faa files
rm -rf *.faa

# Compress the file
bzip2 -9 sc.fas
```

Then we will create the SP proteome file

```
mkdir -p SP
cd SP
wget --quiet ftp://ftp.ncbi.nih.gov/genomes/Fungi/Schizosaccharomyces_pombe_uid127/*.faa
cat *.faa >sp.fas
rm -rf *.faa
bzip2 -9 sp.fas
```

1.2 Download blast

BLAST has new C++ version called blast+. But we will stick with the older version.

Download the BLAST executable from NCBI FTP site:

ftp://ftp.ncbi.nlm.nih.gov/blast/executables/release/2.2.26/

Edit your ~/.bashrc to put the executable on your path. You also may need to create a .ncbirc depending of the version of the BLAST. For version 2.2.26 you really do not need to create one.

1.3 Formatting the BLAST database

For searching a sequence against a database, BLAST requires the database to be formatted in a specific way. There are some pre-formatted databases available from the FTP site:

```
ftp://ftp.ncbi.nlm.nih.gov/blast/db/
```

However, in our case, we need to format our own database. First we will combine the two genomes into one file.

```
bzcat *.bz2 | bzip2 -c >all.fas.bz2
```

The software that formats a fasta file searchable by BLAST is called formatdb. You can view the full list of options by running:

```
formatdb -
```

You can simply run formatdb using the following command

```
formatdb -i <inpufile>
```

This will create a datbase with the same name as the input file. However, this requires a unzipped flat file, which is a no no in our case. You can run formatdb using the following command:

```
bzcat all.fas.bz2 | formatdb -i stdin -o T -n "all"
```

The -o options tells formatdb to create an index of the sequences. This will come handy if we would like to get a sequence out of this database later. -n option is necessary, because when reading from stdin, formatdb has no idea what to call the database.

Depending on your computer ram, you might have to change -v to fit the database entirely in the RAM. For a list of tips and tricks read the formatdb documentation:

ftp://ftp.ncbi.nlm.nih.gov/blast/documents/formatdb.html

1.4 fastacmd

fastacmd is a very useful utility supplied with BLAST. For instruction see:

ftp://ftp.ncbi.nlm.nih.gov/blast/documents/fastacmd.html

1.5 Splitting the query file in small chunks

To run BLAST over cluster we need to split the input fasta file into smaller chunk that will be send over to the cluster for parallel BLAST. We will split the file using a home-grown tool.

```
git clone https://github.com/malaybasu/SeqToolBox.git
```

Now split the fasta file into smaller pieces. The number of piece should approximately equal to the number of nodes that you have in you cluster.

```
echo $(( `bzcat all.fas.bz2| grep \> |wc -l` / 128 ))
perl SeqToolBox/bin/splitfasta.pl -s 86 <(bzcat all.fas.gz)</pre>
```

1.6 Submit jobs to cluster

```
for i in `ls *.fas`
do
qsub -cwd -b y -V -j y -o $i.log -l h_rt=1:00:00,vf=5G \
"blastall -e 0.001 -F T -p blastp -i $i -m 9 -d /scratch/user/malay/all \
>$i.bla && bzip2 $i.bla"
done

# Clean the fas files and log files
rm -rf *.log *.fas

# Create a big file containing all the results
cd ..
bzcat test/*.bz2 | bzip2 -c >all.bla.bz2
```

One important option is -F that switches the compositional filtering on. I suggest that you keep this option on.

1.7 Exercise

Write a software that will parse the all.bla.bz2 file and the individual genome files and find orthologs using the reciprocal best blast algorithm

2 HMMER

HMMER is the only known software for HMM use in bioinformatics. You can download HMMER from http://hmmer.janelia.org/software

There are quite a few software that are bundled with HMMER distribution. But the 4 most common ones are:

- 1. hmmsearch Searches HMMs against protein sequences
- 2. hmmscan Searches protein sequences against HMM library
- 3. hmmbuild Builds HMM from a multiple alignment
- 4. hmmpress Convert a flat file HMM to binary format that can be used with the software

2.1 Exercise

We will search the yeast genome with a profile of P53 gene. Yeast is known not to have P53, but still we will see what we get.

We will first get a bunch of orthologs of P53 from the Homologene database.

```
wget --quiet ftp://ftp.ncbi.nih.gov/pub/HomoloGene/current/homologene.data
```

Now we will grep the file to find p53:

```
cat homologene.data | grep -i tp53 | head -n 10
                                             NP 000537.3
## 460
        9606
                7157
                         TP53
                                 120407068
## 460
        9598
                455214
                        TP53
                                 332847216
                                              XP_001172077.2
## 460
        9544
                716170
                        TP53
                                 114051852
                                             NP 001040616.1
```

```
## 460
                403869
                                              NP_001003210.1
        9615
                        TP53
                                 50978974
## 460
        9913
                281542
                        TP53
                                 28849929
                                             NP_776626.1
                                              NP_112251.2
## 460
        10116
                24842
                         Tp53
                                 189083686
        7955
                30590
                                 425876787
                                              NP_001258749.1
## 460
                         tp53
## 460
        8364
                431679
                         tp53
                                 50054422
                                              NP 001001903.1
## 3959 9606
                7159
                         TP53BP2 112799849
                                              NP_001026855.2
## 3959 9598
                457766 TP53BP2 332812020
                                              XP 003308815.1
```

Looks like cluster 460 contains T53. We will use a bit of perl code to get the accession.

```
cat homologene.data | perl -ane 'print $F[5],"\n" if ($F[0] == 460)'>p53_homologene_ids.txt
```

We will now use NCBI eutils to extract those sesquence. Let's create a fasta file with this ids:

```
for i in `cat p53_homologene_ids.txt`
do
wget -q -0 - "http://eutils.ncbi.nlm.nih.gov/entrez/eutils/efetch.fcgi?\
db=protein&id=$i&rettype=fasta&retmode=text"
done >p53.fas
```

We will first use muscle to align those sequence and create the HMM, then search SC genome with it.

```
muscle -in p53.fas -out p53.aln
hmmbuild --informat afa p53.hmm p53.aln
hmmsearch -o hits.txt ../p53.hmm sc.fas
```

Indeed looks like there is not hit for P53 in yeast.

2.2 Exercize

Search P53 gene in human genome.

3 Problem 1

Reciprocal best blast hit is a way to find orthologs in two or more genome. The algorithm was modified in Inparanoid to find the inparalogs within a genome. Inparalogs are gene that scores higher than the orthologs. Write a python script that will take a all vs all blast result of SC and SP and the individual BLAST file and will report not only the ortholog but also the inparalogs.