Taxonomic classification of SSU rRNA community sequence data using CREST



2014 Workshop on Genomics, Cesky Krumlov

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1. Familiarise yourself with CREST installation

LCAClassifier is the program in the package CREST (Classification Resources for Environmental Sequence Tags) that is responsible for taxonomic classification of sequences based on Megablast alignments to a reference database. On the course lab computers, this program is installed in the directory ~/software/LCAClassifier. The installation directory contains a configuration file that defines the location of database files that the LCAClassifier needs in order to map Megablast results to a taxonomy. Find and open the file lcaclassifier.conf in the LCAClassifier subdirectory parts/etc, either using the graphical or command line interface.

As you can see this configuration file contains references to three different reference databases / taxonomies. By default, these databases are currently available:

- **I) SilvaMod**, based on the SILVA SSURef database (v106), which contains a taxonomically annotated alignment of small subunit (SSU) rRNA sequences from all three domains of life,
- **II) Greengenes**: another SSU rRNA database for bacteria and archaea using an alternative annotation, and
- III) **UNITE**, which instead is a database only for fungi, using the Internal Transcribed Spacer a non-functional rRNA between the SSU and LSU genes.

They are downloaded automatically when LCAClassifier is installed, into the folder *parts/flatdb*. Each database contains the following files, named like the database itself, e.g. for silvamod:

1) silvamod.fasta.n* (.nhr, .nin and .nsq) –binary search index files for BLAST / Megablast. The FASTA-formatted sequences themselves are not needed by BLAST and not downloaded by default, to speed up the installation.

- **2) silvamod.tre** a text file in Newick tree format that defines the topology of the taxonomical tree
- 3) **silvamod.map** a tab-separated text file that specifies a name and rank for each taxon ID in the tree-file

Now, use the command line shell (Terminal window) create a directory for this tutorial under your home directory, called e.g. *CREST_Tutorial*:

```
cd
mkdir CREST_Tutorial
cd CREST Tutorial
```

2. Download the data

In this tutorial we will classify two sets of amplicon sequence data. These sequence sets are called *LC_Prefilter* and *LC_Final* and originate from an alkaline soda lake in Ethiopia ("LC"=Lake Chitu). The difference between the two is that the DNA for the first is collected from a more rough filter and the second results from this filtrate, collected using a more fine filter (so, not a noise-filter but an actual, real filter for water samples). Thus, we expect to only catch planktonic microorganisms in the second dataset.

First, however, we need to download the data using wget.

```
wget
http://services.cbu.uib.no/supplementary/crest/SodaLakeAmplicons.tar.gz
```

Then uncompressed the three files:

```
tar xvzf SodaLakeAmplicons.tar.qz
```

As you can see, the folder contains several files, including: *LC_Prefilter_F_Good.fa*, *LC_Final_F_Good.fa* and *LC.sff*. The first two are FASTA-files, de-noised and annotated using AmpliconNoise to represent only unique sequences with different numbers of reads. The second is in SFF-format (standard flowgram format), which is the raw data from 454 pyrosequencing. If you want, and if the service is fast enough, you can

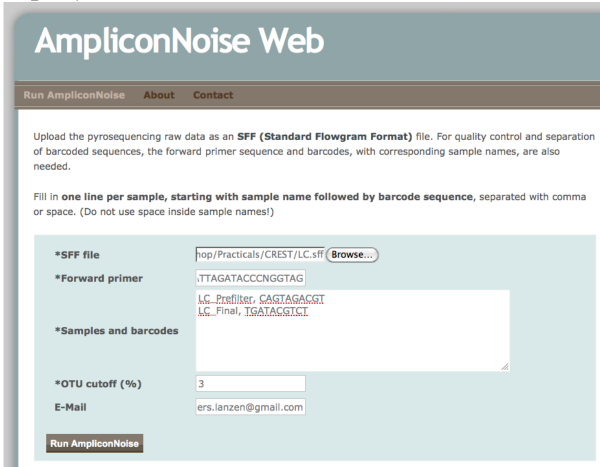
submit this SFF-file to the AmpliconNoise web server (see below). Otherwise:

- 1) use the uncompressed files *AN_stats.txt* and *OTU_Diversity_estimates.txt*, which are already included, or,
- 2) if you are not interested in denoising and diversity estimates, skip the next step and move directly to step 4

3. De-noising the pyrosequencing raw data using AmpliconNoise (optional)

If this tutorial is run in a large group, not everybody can do this at once. If you wish, proceed to step 4 and come back to this step later.

Open a web browser and go to http://apps.cbu.uib.no/ampliconnoise. Upload the SFF file and copy / paste the following sequence into the field "Forward primer": ATTAGATACCCNGGTAG and the following to the field "Samples and barcodes": "LC_Prefilter, CAGTAGACGT" (enter) and "LC_Final, TGATACGTCT". See the screenshot below



Click Run and wait.

When the job is finished, download and look at the files *AN_Stats.txt* and *OTU_Diversity_estimates.txt*. (Note that these are already included in the tutorial dataset, for the inpatient!)

- How many raw reads did you submit in the SFF file from each dataset?
- How many reads remain after filtering and chimera removal?
- How many unique sequences and 3% OTUs do the remaining reads constitute?
- What is the Shannon diversity index?

Download the files *OTU_All_Samples.fa* and *OTU_table.csv* for later. Also download the file *Clean_Sequences_w_Abundance.tar.gz* and uncompress it. This is your de-noised sequences (the same as those provided already, in case of skipping this step). You can see that each sequence name ends with an underscore and a number, e.g. "_3". This indicates the number of reads representing a unique sequence and can be understood by the LCAClassifier later on.

4. Aligning files to the reference database (SilvaMod) using Megablast

The first step in the CREST workflow is alignment to a reference database, using Megablast, which is a faster version of BLAST (Basic Local Alignment Search Tool) for nucleotide sequence alignments. We will use the **SilvaMod** database made from SILVA's SSURef alignment of full-length SSU rRNA sequences, release 106. Using the resulting alignments (BLAST results), taxonomic classification is then done with the **LCAClassifier** program. The LCAClassifier only supports results from the NCBI *blastall* implementation of Megablast, in XML format. It does not work with results from the newer BLAST+ implementation.

Align the sequences in *LC_Final_F_Good.fa* to SilvaMod using:

```
megablast -i LC_Prefilter_F_Good.fa -b 50 -v 50 -m 7 -d
~/software/LCAClassifier/parts/flatdb/silvamod/silvamod.fasta -a 2 -o
LC Prefilter silvamod.xml
```

This will take some 5 minutes. Meanwhile, have a look at the the arguments you gave to the megablast command and what they mean:

- -i: the nucleotide sequence input file
- -b 50 -v 50 : Report only the 50 best alignments for each sequence
- -m 7: Produce output in XML format
- -d: the reference database to use*.
- -a 2: Use two CPU cores / threads.
- -o: destination file of the Megablast output

If you don't get any error message, the Megablast alignment has probably worked fine. You can also have a look at the resulting file in a text editor, to familiarise yourself with the BLAST XML format, which is the same for Megablast and normal BLAST. However, it is not really intended for being read by human beings and is a bit too structured for us.

^{*}Note that the file *silvamod.fasta* itself is actually missing by default, but is not needed since MEGABLAST only uses the search index files produced by the command *formatdb* from *silvamod.fasta*.

Then, align the other FASTA-file:

```
megablast -i LC_Final_F_Good.fa -b 50 -v 50 -m 7 -d
~/crest/LCAClassifier/parts/flatdb/silvamod/silvamod.fasta -a 2 -o
LC Final silvamod.xml
```

Again, this takes a few minutes through the QIIME virtual box. Have patience, go for some coffee and relax.

5. Taxonomic classification using LCAClassifier

Classification with the LCAClassifier program using default parameters is quite easy and carried out using the command **classify.** To classify the Megablast-aligned *LC_Prefilter* dataset, simply type:

```
classify LC_Prefilter_silvamod.xml
```

This will result in two files. The first, named

LC_Prefilter_silvamod_Composition.txt, is a tab-separated text file, listing the number of reads, relative abundance, unique sequences and a Chaoestimate of minimum diversity for each taxon, at different rank levels (domain, phylum, class, order, family and genus). The number of reads classified at each rank level is also summarised. Open this file in a spreadsheet editor (like LibreOffice; *make sure to chose only "tab" under "separated by"*)

- What proportion of the original reads could be classified to at least family level?
- What is the relative abundance of *Arthrospira* (a common alkaliphilic cyanobacteria)? How many unique sequences are represented in this genus?
- How many unique sequences were assigned to the family Rhodobacteraceae and what is the Chao-estimate for total number of unique sequences?

- How many eukaryotic sequences are there? Can you figure out why, although the primers used target 16S only?

The other file, *LC_Prefilter_silvamod_Tree.txt*, lists the number of reads assigned in a simple space-delimited tree-format. Here are the first ten lines of the file:

```
head LC_Prefilter_silvamod_Tree.txt

root: 561
Cellular organisms: 561
Archaea: 27
Euryarchaeota: 27
Methanomicrobia: 9
Methanomicrobiales: 6
Methanocalculus: 6
Methanosarcinales: 3
Methanosaetaceae: 2
Methanosaeta: 2
```

6. Comparison of composition between datasets

The LCAClassifier can also classify several Megablast result files at the same time, providing that they were aligned to the same reference database. First delete or rename the old *LC_Prefilter_Silvamod_Composition*.txt file. Then, try this out:

```
classify -o *.xml
```

This results in the same two types of result files for each dataset (*LC_Prefilter* and *LC_Final*). However, empty assignments are inserted with zero abundance for each taxon present in at least one dataset, where no assignments to the corresponding taxon was made in a particular dataset. This helps to compare datasets. The option –o instructs the LCAClassifier to also write results in alternative output format (*All_Composition.txt*).

Open LC_Prefilter_Silvamod_Composition.txt and LC_Final_silvamod_Composition.txt in a spreadsheet editor (e.g. LibreOffice). Libreoffice->Spreadsheet->Insert->Sheet form file->Separated by tab

Copy the last four columns ("Abundance" to "Chao") of one file into the right of the columns of the other one, so that each taxon is compared for the two datasets side by side.

- Can you find any taxa overrepresented in the prefilter sample, with an average abundance > 1% and less than half in the "final" sample? These are likely to be bulky, filamentous or non-planktonic organisms.
- How about a dominant taxon with almost twice the abundance in the *LC_Final* dataset?
- What is the most diverse genus, in terms of number of unique denoised sequences?
- Have a look at the file *All_Composition.txt*. Can you figure out how many sequences that were assigned to the domain Bacteria but could not be assigned to any particular phylum.

7. Extracting sequences from a taxon

The LCAClassifier can also write output files that specify the assignment for each individual sequence. Using option **-p** (or **--rdp**), the identifier of each sequence and its predicted taxonomical path is written with suffix "_Assignments.txt". For example:

SedimentX_47_5 Cellular organisms;Archaea;Euryarchaeota; Methanomicrobia;ANME-1;ANME-1a4

Another useful option is to write a new FASTA file with taxonomic annotations added in the FASTA header. This is done using option -a (or -fasta). By default, only the aligned part of each sequence is written and entries that could not be aligned are omitted. Alternatively, the entire sequences can be written. The file to read in sequences from then must be specified using -i (or --fastain).

We will now produce these files for our representative OTU sequences (Operational Taxonomic Units, basically clusters of similar, unique sequences). Use the file *OTU_All_Samples.fa*, which contains representative sequences for each OTU in the two datasets:

```
megablast -i OTUs_All_Samples.fasta -d
~/software/LCAClassifier/parts/flatdb/silvamod/silvamod.fasta -b 50 -v
50 -m7 -a 2 -o OTUs_silvamod.xml
classify -i OTUs All Samples.fasta -a -p OTUs silvamod.xml
```

This produces two files with prefix ".fasta" and ".txt" as described above. The first can be used for extracting all sequences from a particular taxon, using the LINUX command **grep.** For example, to save all archaeal sequences to the file *A.fa*:

```
grep Archaea OTUs_silvamod_Assignments.fasta -A1 > A.fa
```

- How many archaea are there?

```
(Hint: grep -c '>' A.fa, Note that the single quotes are important!)
```

Some of the sequences are assigned to "Unknown" taxa. This means that the minimum sequence similarity filter of the LCAClassifer has prevented it from being classified to a higher rank. These sequences may be interesting because of their novelty. Have a look at these sequences:

```
grep Unknown OTUs silvamod Assignments.fasta -A1 > Unknown.fa
```

The second last sequence ("C56") is assigned to an unknown order (under *Alphaproteobacteria*). This means that it is less than 90% similar to the closest reference sequence and thus cannot be assigned to a particular order, but instead to class *Alphaproteobacteria*. Copy the

sequence to NCBI Blast (http://www.ncbi.nlm.nih.gov/blast) and have a look at the closest sequence matches. Note from what type of environment the most similar sequence hit is.

To read an overview of all options available with the LCAClassifier, type:

```
classify --help
```

8. Classification using Greengenes

As an alternative to the SilvaMod Taxonomy, the LCAClassifier can also be used together with the Greengenes Taxonomy. Similarly to SILVA's (and "SilvaMod"), this is basically a reference database consisting of environmental sequences and type strains annotated taxonomically based on the clustering, or in other words the topology of the distance tree. The difference from SILVA is that the annotations in Greengenes were made automatically, using a heuristic algorithm. To read more about the Greengenes taxonomy, see McDonald et al (2012), 'An improved Greengenes taxonomy with explicit ranks for ecological and evolutionary analyses of bacteria and archaea', ISME I, 6:610-618.

As the reference database is different, the first step is to align the sequences to the Greengenes reference database, which can be found in ~/crest/LCAClassifier/parts/flatdb/greengenes. To align e.g. the LC_Final sequences:

```
megablast -i LC_Final_F_Good.fa -b 50 -v 50 -m 7 -d \sim/crest/LCAClassifier/parts/flatdb/greengenes/greengenes.f asta -a 2 -o LC_Final_GG.xml
```

To classify the sequences based on the Megablast result, you need to tell the *classify* command to use the Greengenes taxonomy, since SilvaMod is the default.

```
classify -d greengenes -p LC Final GG.xml
```

Open *LC_Final_GG_Composition.txt* in a spreadsheet editor.

- What proportion of reads could be classified at **genus level**?

 Compare it to the community composition based on the SilvaMod taxonomy. What major differences can you find between the results based o the two reference databases / taxonomies, on family level?

9. Classification with MEGAN (optional)

In addition to the LCAClassifier, the program MEGAN also supports classification using the CREST scheme. MEGAN has a graphical user interface and many functions for taxonomic and functional classification of metagenomic sequence data. We will revisit these functions in another practical of this course.

By default, MEGAN uses the NCBI taxonomy for assignment. To change this to the SilvaMod taxonomy, select "Use Alternative Taxonomy..." under Edit>Preferences. Then navigate to the directory ~/software/LCAClassifier/parts/flatdb/silvamod and select the file silvamod.tre. If everything works correctly, the SilvaMod taxonomy should now be enabled and you should see the following output in the Messages window:

Now, import the BLAST results, using File>Import from BLAST. Select one of the Megablast XML files, then go to the sheet LCA Params and change the settings to the following: Min support=1, Min score=155 and Top Percent=2. Also, tick the box "Use Percent Identity Filters". Then, click Apply!

An interactive taxonomy tree will now appear, listing the number of assignments to different taxa and also symbolising the abundance of a taxon by a circle, whose area is proportional to the number of assignments. Try to explore the tree by collapsing and uncollapsing nodes and by right-clicking a node and selecting Inspect.