#### A Guide to the stand-alone version of MLTreeMap:

MLTreeMap is a software framework, designed for phylogenetic and functional analysis of metagenomic data. It searches for instances of marker genes on nucleotide sequences and deduces their most likely origin in a set of reference phylogenies. The current version of MLTreeMap can be downloaded and installed individually. MLTreeMap runs on Mac and Linux systems.

#### A) Installation of MLTreeMap

## Step 1

Unzip and unpack the file MLTreeMap\_package\_2\_04.tar.gz.

## Step 2

Enter the directory MLTreeMap package 2 04/install/.

Type make.

The make program now creates the data structure of MLTreeMap (to be found in MLTreeMap\_package\_2\_04/mltreemap\_2\_04/), and compiles most needed sub-programs (hmmalign, Genewise and RAxML).

#### Step 3

BLAST and Gblocks have to be added manually. For this enter the directory MLTreeMap\_package\_2\_04/install/sources/BLAST/. Here you will find a collection of BLAST binaries. Choose the one appropriate for your system and copy it to the directory MLTreeMap\_package\_2\_04/mltreemap\_2\_04/sub\_binaries/.

```
Repeat this for Gblocks (to be found in MLTreeMap_package_2_04/install/sources/Gblocks/).
```

MLTreeMap is now ready to use. You can copy the directory mltreemap\_2\_04 to any place you like (as well as renaming it). The only dependency of MLTreeMap is that you must have Perl installed on your system.

### B) Usage of MLTreeMap.

MLTreeMap has to be accessed on the command line.

```
An example for a valid input command is:
./mltreemap.pl -i example_input/rubisco.txt
```

This will analyze the sequence in the file rubisco.txt and write the result to the output directory mltreemap 2 04/output/.

Further (optional) input parameters are:

- -b number of bootstrap replicates (Default 0 i.e. no bootstrapping).
- -c usage a computer cluster (0 = no cluster (default), s = sun grid).
- -f RAxML algorithm (v = Maximum Likelihood (default), p = Maximum Parsimony).
- -g minimal sequence length after Gblocks (default = 50).
- -l long input files will be split into files of n sequences each (default 2000).
- -o output directory (default: output/).
- -s minimum bitscore for the blast hits (default 60).

-t phylogenetic reference tree (p = MLTreeMap reference tree (default), g = GEBA reference tree).

## C) The MLTreeMap output.

MLTreeMap searches for phylogenetic and functional marker genes. As soon as the results can be assigned to a specific marker gene, they are labelled accordingly by the first character of the output files.

# Phylogenetic analysis:

- a 16s rRNA reference data and tree
- b 18s rRNA reference data and tree
- g GEBA reference tree
- p MLTreeMap reference tree

## Functional analysis:

- c Photolyase & cryptochrome
- d Reverse dissimilatory sulfite reductase (DsrAB)
- h NifH (K02588)
- m Methane & ammonia monooxygenase
- n NifD (K02586)
- r RuBisCo (COG1850)

Additionally, each output file after the RAxML step gets a header line, providing this information in words. In case of the RuBisCO example from above, the final output file would look as follows:

# Functional analysis, RuBisCO:

Placement weight 100%: Assignment of query to Acry 1067 (Acidiphilium cryptum) (6).