OMA stand-alone

CBRG, ETHZ

Contents

1	Introduction				
2	Downloads				
3	Installation				
4	Usage 4.1 Parallelization 4.1.1 Parallelization with LSF, PBS Pro, Slurm, or SunGridEngine 4.1.2 LSF 4.1.3 Sun Grid Engine (aka Oracle Grid Engine) 4.1.4 PBS Pro 4.1.5 Slurm 4.2 Required Resources 4.3 Adding/Updating new genomes				
5	File Formats 5.1 Input Files 5.2 Output Files 5.2.1 OMA Output 5.2.2 ESPRIT Output				
6	Parameters				
7	Getting help				
8	License				

1 Introduction

You can run OMA as a stand-alone program. Included are the algorithms for OMA itself plus its addition ESPRIT. The software can be installed on Linux (x86, both 64bit and 32bit) and MacOSX (x86, both 32bit and 64bit).

For a short summary and pointers to detailed algorithmic publications of OMA, please refer to the OMA browser page:

http://omabrowser.org/oma/about/

For background info on ESPRIT, please refer to this article:

http://dx.doi.org/10.1093/bib/bbr038

If you have specific questions about the installation or the usage of OMA, please contact adrian.altenhoff at inf.ethz.ch or christophe.dessimoz at unil.ch.

2 Downloads

The current version of OMA stand-alone can be found here: OMA.1.0.5.tgz

3 Installation

You do not need to install OMA stand-alone on your system; the script will also run if you just call it by using the complete path to bin/oma in the installer folder. But we still encourage you to run the installer script, since it makes working with OMA considerably more convenient.

To install OMA stand-alone on your system, download the installer, untar the package and run the included installer script:

```
curl http://omabrowser.org/standalone/OMA.1.0.5.tgz -o oma.tgz
tar xvzf oma.tgz
cd OMA.1.0.5
./install.sh /your/install/prefix
```

If you do not choose an install prefix, OMA will be installed in /usr/local/OMA (for this, you might need to install it using the root account or sudo).

After installation, make sure the bin folder of OMA is in your PATH variable, e.g., if you are using bash and used /your/install/prefix as installer prefix, add a line in /.profile such as:

```
export PATH=\$PATH:/your/install/prefix/OMA/bin
```

For other shells, choose the appropriate syntax.

4 Usage

First, set up a working directory. Copy the file parameters.drw into this folder and change it to your needs. Create a directory DB in your working directory that holds the genome data in FASTA format (see "File formats") and copy your data into this directory. If you want to use ESPRIT, the FASTA file containing the contigs should be called {YourGenome}.contig.fa. Then, simply call OMA from your working directory to run OMA and/or ESPRIT

If you have not installed OMA yet, use the complete path to bin/oma in the installer folder to start the script.

As an example, assume you installed OMA in /your/install/prefix and want to use ES-PRIT on two genome files and one file with contigs (all in /home/you/fasta, do something like this:

```
# create working directory
mkdir myWorkingDir
cd myWorkingDir
# create DB directory in working directory
mkdir DB
# copy FASTA files into DB directory
cp /home/you/fasta/yourFirstGenomeFile.fa DB/
cp /home/you/fasta/yourSecondGenomeFile.fa DB/
cp /home/you/fasta/yourContigFile.contig.fa DB/
cp /your/install/prefix/OMA/OMA.1.0.5/parameters.drw ./
# adjust parameters
vim parameters.drw
# run OMA
OMA
```

To get a first impression of OMA you could cd into the ToyExample directory, have a look at parameters.drw and run OMA to process our example files.

4.1 Parallelization

The all-against-all phase of OMA is the most time-consuming one, but it can be parallelized (unlike all other steps, which cannot run in parallel). The way it works is that the parameter "AlignBatchSize" and the total number of genomes (n) will determine into how many chunks the all-against-all phase is divided. AlignBatchSize will split the n*(n-1)/2 genome pairs further into chunks of at most "AlignBatchSize" alignments. The larger AlignBatchSize is, the more and smaller jobs will be executed.

Scheduling is straightforward: all compute processes need to start from the same directory, and each one will try to do an equal amount of chunks sequentially. However, before starting a new chunk, each process ensures that it has not yet already been processed by another process (i.e. no result file yet exists).

On a single computer with multiple processors and/or cores, it is recomended to start "N" parallel processes with the "-n" option, i.e.

```
OMA -n 5
```

will start 5 parallel jobs. Note that on HPC with schedulers there is a better way described below 4.1.1

Therefore, there is not need to specify which parts are to be done by which process. One should only ensure that all processes start from a shared directory, such that each chunk gets computed by a single process only.

4.1.1 Parallelization with LSF, PBS Pro, Slurm, or SunGridEngine

With a scheduler such as LSF or SGE, running parallel jobs is particularly easy, as the parallel jobs can be start using as a job array. OMA will automatically spread the work for the allagainst-all among all processes. For a brief discussion on the required resources refer to the section below4.2 Do not start OMA with the -n option for that. Instead, use a job array with one of the supported schedulers below (the example is to start 100 jobs in parallel)

4.1.2 LSF

```
bsub -J "OMA[1-100]" -o "out.%I" "OMA"
```

4.1.3 Sun Grid Engine (aka Oracle Grid Engine)

```
qsub -t 1-100 "OMA"
```

4.1.4 PBS Pro

Prepare a job script called e.g. job.sh:

```
#!/bin/bash
# set the number of nodes and processes per node
#PBS -l select=1:ncpus=1:mem=1000mb
# set max wallclock time
#PBS -l walltime=01:00:00
#PBS -J 1-100
OMA
```

The script can then be submitted as follows:

```
qsub job.sh
```

4.1.5 Slurm

```
sbatch --array=1-100 -N1 <<EOF
#!/bin/sh
/absolute/path/to/bin/OMA
EOF</pre>
```

4.2 Required Resources

Depending on the dataset to be analysed, OMA can require quite a significant amount of computational resources, i.e. RAM and cpu time. Most computing time is spent to compute the all-against-all sequence alignments, which is why this part has been parallelized. Although cpu intensive this phase does not require too much memory. As a rule of thumb you can assume it requires roughly 10 times the size of your DB folder, but at minimum 650MB. The second part of OMA runs sequential on a single core, but it requires a lot more memory: Again, as a rule of thumb, count on 400MB * pow(nr_genomes, 1.4). Obviously this depends also a lot on the size of your genomes. 60 metazoas have been successfully computed using 120GB, and for the same number of bacterial genomes, 50GB were reported to be enough.

Because of this imbalance regarding the required memory the OMA starter script has command line option –s to run only the first part of the OMA pipeline. Using this option, the computation can be split into a parallel phase with little memory requirement , and a single process requiring a lot more RAM. Running the OMA thus becomes a two stage process like this:

```
# first stage, little memory:
OMA -s
# once first stage terminated, run second stage
OMA
```

Note that this staging is mostly useful on computing clusters where often memory and cpu time has to be reserved at job submission.

4.3 Adding/Updating new genomes

It is possible to add new genomes without recomputing the all-against-all phase for pre-existing genomes. To do so, simply add the new fasta databases in the DB/ directory and re-run OMA. Likewise, it is possible to update a genome by deleting the old genome from the DB/ directory, and adding a new file. **Important:** to avoid clashes with previously computed results, the updated genome must use a different filename than the any previously computed result.

5 File Formats

5.1 Input Files

OMA uses two different input formats: FASTA files for genome input and a Darwin file for parameter input.

The Fasta format is explained in detail on wikipedia.

As almost everywhere else, OMA uses the greater-than symbol ">" to distinguish labels from sequences. Each sequence in a genome is supposed to have its own label. Have a look

at the FASTA files included in ToyExample/DB in our installer package for some example files.

In case your genomes contain multiple alternative splicing variants, you can add a text-based file per genome calledDB/{YourGenome}.splicethat put together the different splicing variants, e.g. to indicate that the three splicing variants ENSP00000384207, ENSP00000263741 and ENSP00000353094 are encoded by the same gene, add the following line to the splice file:

ENSP00000384207; ENSP00000263741; ENSP00000353094

OMA requires that the individual IDs are uniq prefixes of your FASTA headers.

If you want to use ESPRIT, make sure that FASTA files containing contigs are called {YourGenome}.contig.fa. So if you want to experiment with some mouse genome, call the FASTA file mouse.contig.fa or mymouse.contig.fa or something similar.

Parameter files use Darwin syntax. Key-value-pairs are written as

key := value;

Note the colon in := and the semicolon at the end of the line. If your parameter file does not use valid Darwin syntax, OMA will print out a short message and stop its execution.

5.2 Output Files

5.2.1 OMA Output

The output of OMA gets written to files stored in a directory Output in your working directory. This can be changed by changing the OutputFolder parameter, There are text files and directories organized as described in Table 1.

5.2.2 ESPRIT Output

ESPRIT stores its output files in a directory called EspritOutput in your working directory. The output consists of three text files and one tarball. In the tarball, FASTA files with the MSAs of the hits ESPRIT found are stored. The other three files are explained in detail in Table 2.

6 Parameters

All parameters for OMA and/or ESPRIT are set in a parameters file. There is an example file in the OMA installer package; we encourage you to copy this file into your working directory and change it to your needs.

The parameter file consists of two main parts: First, general parameters for OMA are set; see Table 3 for detailed explanations. Second, more specific parameters that only affect the ESPRIT algorithm can be changed. These parameters are explained in Table 4. Note that changing the ESPRIT parameters will not have an effect unless you set the boolean variable UseEsprit to true.

Filename or Directory	Contents
Map-SeqNum-ID.txt	Lists all genes of all datasets with their unique sequence number and the labels read from the FASTA files.
OrthologousGroups.txt	The groups of orthologs are given as one per row, starting with a unique group identifier, followed by all group members, all separated by tabs.
OrthologousMatrix.txt	More compact version of OrthologousGroups.txt. The groups of orthologs are given as matrix with group per row and one genome per tab-separated column. Numbers refer to entry number as listed in the file Map-SeqNum-ID.txt.
${\tt OrthologousGroups.orthoxml}$	The OMA groups of orthologs stored in orthoxml format.
OrthologousGroupsFasta/	Each OMA group is provided as a separate Fasta file, with the species name as identifer. This format is particularly suitable as starting point for a phylogenetic reconstruction.
PairwiseOrthologs/	The textfiles in Output/PairwiseOrthologs are named according to {genome a}-{genome b}.txt and consist of a list of pairwise orthologs for the two given genomes. Every pair is listed only once, and in no particular order. Each line in the file contains one pair; all fields are separated by tabs. In the first two field, the unique IDs of the proteins are given. The next two fields contain the labels of the proteins, and in the last two fields, the type of orthology and (if any) the OMA group is given.
OrthologousPairs.orthoxml	The pairwise orthologs stored in orthoxml format. Each group in the file will have orthologs between genes from only two genomes.
HierarchicalGroups.orthoxml	The hierarchical groups of orthologs in OrthoXML format. A detailed description of how these groups are computed is forthcoming.
EstimatedSpeciesTree.nwk	The inferred species tree on which the hierarchical groups inference procedure is based, in Newick format.
HOGFasta/	For each toplevel HOG group we provide a separate Fasta file with all protein sequences clustered to it. This format is particularly suitable as starting point for a phylogenetic reconstruction of a gene tree.
used_splicing_variants.txt	If UseOnlyOneSplicingVariantis activated and splicing information is available, the variant which has been used for calling the orthologous relations is stored in this file. The format is a tab-delimitted text file with the species in the first column and the id of the used splicing variant in the second column.

Table 1: Contents of the OMA output files

Filename	Contents	
params.txt	This file is kept as a reference and contains all parameters used in the	
	current run.	
hits.txt dubious.txt	current run. All hits found by ESPRIT are listed in this file. It is a list of contiguordered according to their position relative to the putative ortholog. Ea line describes one contig, the fields are separated by tabs. In the first file the fragment pair ID is printed; the next two fields contain the labels the first and second fragments found in this hit. The forth and fifth file contain the label of the corresponding full gene and its genome name. The follows the distance difference between the two fragments and the number of positions between them (i.e. the gap); at last, an array is listed containing the IDs of all s3 genes corresponding to this hit.	
	which triplets have been filtered out, have a look at dubious.txt where they are still listed. The file format is the same as for hits.txt.	

Table 2: Contents of the ESPRIT output files

7 Getting help

The preferred way to get help about OMA is via the Biostars community resource. Please consider asking your question there, including the tags "OMA" and "orthologs". If your question requires privacy, we are also reachable by email at contact@omabrowser.org.

8 License

OMA standalone is licensed under the Mozilla Public License Version 2.0. For more info, please consult the following page:

http://www.mozilla.org/MPL/2.0/

In a nutshell, OMA standalone is open source and free for commercial and non-commercial

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Parameter	Meaning	Default
InputDataType	Type of input sequences. This can be set either to 'AA' for amino acid sequences or 'DNA' for nucleotide sequences	AA
OutputFolder	Folder to which the output is written. At each run, the content of this folder will be overwritten. Don't store any important files in it. The OutputFoldermust not contain any spaces.	Output
ReuseCachedResults	If you want to recompute everything from scratch everytime the script is run, set this to false.	true
AlignBatchSize	In the all-against-all phase, each genome pair is split in smaller chunks of AlignBatchSizeprotein comparisons. The larger this number, the longer each unit runs, and the fewer files get produced. This allows to adjust the frequency of milestone steps (e.g. in case of computer crash) or to process few but large genomes with many CPUs efficiently.	1000000
MinScore	Alignments which have a score lower than MinScore will not be considered. The scores are in Gonnet PAM matrices units.	181
LengthTol	Length tolerance ratio. If the length of the effective alignment is less than LengthTol * min(length(s1), length(s2)), then the alignment is not considered.	0.61
StablePairTol	During the stable pair formation, if a pair has a distance provable higher than another pair (i.e. StablePairTol standard deviations away) then it is discarded.	1.81
VerifiedPairTol	Tolerance parameter for the detection of differential gene losses using a third genome. The larger the tolerance, the liberaler the algorithm assigns orthologous relations. A detailed description is provided here.	1.53
MinSeqLen	Any sequence which is less than MinSeqLen amino acids long in regular genomes is not considered.	50
UseOnlyOneSplicingVariant	Enables/disables the filtering on a single representative splicing variant. If enabled, OMA selects the variant that has the most homologous matches with all other genomes. Orthology inference is then only based on this variant. If disabled, alternative splicing variants will usually be infered as paralogs.	true
StableIdsForGroups	Enables/disables the generation of stable identifiers for OMA groups (and Hierarchical Groups if their computation enabled). The identifier consists of a prefix to determine the type of the group ('OMA' or 'HOG'), and a subsequence of the amino acid sequence uniquely present in this group. The computation of these ids might require a substantial amount of time. The ids are stored in the OrthoXML files only.	false
DoHierarchicalGroups	Enables/disables the computation of the hierarchical groups. This steps requires substantial computating power.	true
MaxTimePerLevel	Define maximum amount of time (in sec) spent by the program for breaking every connected component of the orthology graph at its weakest link on a given taxonomic level. If set to a negative value, no timelimit is enforced.	1200
SpeciesTree	The hierarchical groups require a (partially) resolved species phylogeny. With the parameter SpeciesTree the user can specify a phylogeny in Newick-format, or,	estimate

Parameter	Meaning	Default
UseEsprit	You can either set this to true, which will enable ES-	false
	PRIT and shut down the parts of OMA that are not	
	directly needed for ESPRIT, or set it to false to make	
	no use of ESPRIT at all.	
DistConfLevel	Confidence level variable for contigs. This is the param-	2
	eter tol described in the paper.	
${\tt MinProbContig}$	Minimal proportion of genomes with which contigs form	0.4
	many:1 BestMatches to consider that we might be deal-	
	ing with fragments of the same gene. This is the param-	
	eter MinRefGenomes described in the paper, normalized	
	by the total number of reference genomes.	
${\tt MaxContigOverlap}$	Maximum overlap between fragments of same gene from	5
	different contigs.	
${\tt MinSeqLenContig}$	Any sequence which is less than MinSeqLenContig	20
	amino acids long in contigs is not considered.	
MinBestScore	Minimum best score for BestMatch in scaffold recogni-	250
	tion.	

Table 4: ESPRIT parameters