

# **Moa Documentation**

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# **CONTENTS**

1	Quic	k links	3		
2	Table of contents:				
	2.1	Goals	5		
	2.2	Introduction	5		
	2.3	Installation	7		
	2.4	Three core templates	10		
	2.5	Synchronizing jobs	11		
	2.6	How to write a template	12		
	2.7	Command reference	15		
	2.8	Templates	25		
	2.9	Moa API	47		
3	More	More information			
4	4 Indices and tables				
Рy	thon N	Module Index 1	65		
In	dex	10	67		

Lightweight, command line, workflows for bioinformatics

Moa aims to assist a bioinformatician to organize, document, share, execute and repeat workflows in a command line environment without losing any of the flexibility of the command line, and, at all times giving the user full access to all aspects of the workflow (see also *Goals*).

\*\*NOTE: both the software and the manual are under development. Things might change.\*\*

CONTENTS 1

2 CONTENTS

## **CHAPTER**

## ONE

# **QUICK LINKS**

- Source code: https://github.com/mfiers/Moa
- Issue tracker: https://github.com/mfiers/Moa/issues
- Old issue tracker <a href="http://moamoa.lighthouseapp.com/projects/73665-moa/overview">http://moamoa.lighthouseapp.com/projects/73665-moa/overview</a>
- Python Package Index: http://pypi.python.org/pypi/moa/ (note this is not updated regularly might not work)
- Source code at Github
- PDF version of the manual

# **TABLE OF CONTENTS:**

## 2.1 Goals

Moa has as objective to assist in keeping a bioinformatics project:

#### • Organized:

Moa facilitates project organization in many (smaller and more major) ways, for example by providing a uniform way to capture commands as Moa jobs. Each Moa job is linked to a specific directory, and has all configuration, templates, data, and intermediate data available as files in this directory structure.

#### • Documented:

Moa provides the possibility to add a title, description and changelogs to each job.

## • Reproducible

By having all templates and configuration copied into a workflow - the workflow does never change (unless the user wants it to), even if templates in the repository change. Moreover, all templates are easy to find & inspect so it is always clear what happened.

## • Reusable & Shareable:

Moa provides reusable templates. New templates are easy to create, adapt and share. Workflows can be archived and reused with different data.

#### • Flexible:

Moa provides a good number of hooks to insert custom code into a workflow, making that code part of the workflow. This ensures maximum flexibility.

## 2.2 Introduction

These days, generating massive amounts of data is an everyday element of biological research; and almost all projects have a computational biology, or bioinformatics, components. Such embedded work commonly consists of chaining a number of 3<sup>rd</sup> party tools together, often with some data manipulation in between the steps. It is important to have such projects properly organized, particularly when a projects grows bigger.

There are many different ways to organize bioinformatics projects. Many bioinformaticians use the command line or tailor made scripts to organize and automate their work. This approach has obvi-

ous advantages, most importantly flexibility. Potential downsides to scripting are that a project easily becomes disorganized and untraceable unless measures are taken.

*Moa* aims to assist in organizing, automating and maintaining a command line bioinformatics project without loss of flexibility.

## 2.2.1 Example

The best way to understand how Moa can help you to achieve this is by an example. A Moa workflow consists of separate Moa jobs. A workflow is typically organised as a directory tree, where the structure of the tree reflects the structure of the project. So, Starting a Moa project starts with outlining a directory structure to contain the workflow:

```
$ mkdir test.project && cd test.project
$ mkdir 00.proteins

( copy or link some protein sequences into 00.proteins )
$ mkdir 10.blast
$ cd 10.blast
```

An important feature of Moa is that each separate analysis step is contained within a separate directory. Two Moa jobs never share a directory. This forces a Moa user to break a workflow down to atomic parts, which is typically beneficial to the organization and coherence of a workflow. The order of steps is easily ordered by prefixing directory names with a number. Note that these prefixes are not enforced by Moa; any alphabetical organization would work as well. Once a directory is created, a Moa job can be created:

```
$ moa new blast -t "demo run"
```

All interaction with Moa is done through a single command: *moa*. It is, at all times, possible to get help on the use of the *moa* command by invoking *moa* –*help*. The command above creates a *BLAST* job titled "demo run" in the current directory. All Moa related files are stored in a (hidden) sub-directory names .*moa* (have a look!). A Moa job consists, amongst others, of a configuration file and a number of template files. All template files are copied into the .*moa* directory. This ensures that a workflow remains the same over time, even if the templates are updated (*moa refresh* would update a template to the latest version).

Another topic in which Moa tries to help is by embedding (some) documentation. In the above command line the -t parameter sets a mandatory project title (a job won't execute without a title).

Obviously, telling a Moa job to do a BLAST analysis is not enough, some extra information will need to be given:

```
$ moa set db=/data/blast/db/nr
```

A few things could be noted here. Important is that you do not use spaces around the = sign. If you want to define a parameter with spaces, use quotes (*key="value with spaces"*), and be aware of bash interpretation. A safe way of entering complex parameters is by running *moa set db* and Moa will query you the value.

Another point is that Moa does not give you a response. You can check the current job configuration using *moa show*, which would at this moment result in something resembling:

```
db L /data/blast/db/nr
input E (undefined)
jobid L blast
title L demo run
```

Note the variable *db* and *title*, which were set earlier. If you run *show -a*, more parameters will be revealed, amongst which is *program*. We will now set two more variables:

```
$ moa set program=blastp
$ moa set input=../00.proteins/*.fasta
```

The last statement defines the input files to blast. Once all is set you can actually run the BLAST analysis with:

```
$ moa run
```

Now Moa performs the BLAST analysis on the input files. The output can be found in the *out* sub-directory. As an extra, the Moa *blast* template generates a *blast\_report* file with simple one line report for the best five hits of each query sequence. If you, for example, would like to check for the presence of dicer genes in your query set, you could *grep* this file:

```
$ grep -i dicer blast_report
```

Command line operation of data files can be very powerful, and this would be a typical operation for a command line bioinformatician. Moa lets you capture this and thus make it a part of the pipeline. Try:

```
$ moa set postcommand
```

and, at the prompt enter:

```
postcommand:
> grep -i dicer blast_report > dicer.out
```

If you now rerun *moa*, the BLAST job will not be repeated, but the *postcommand* will be executed and a *dicer.out* file will be generated. (note, there is also a *precommand*)

## 2.3 Installation

## 2.3.1 Prerequisites

Moa is developed and tested on Ubuntu and RHEL and is expected to operate without much problems on all modern Linux distributions. Moa has the following prerequisites (and a large number more for all templates). Version numbers are an indication, not strict prerequisites. Other, even older, versions might work.

- Python (2.6 or 2.7). Moa will not work with versions earlier, or with 3.0 and up
- Git (1.6). Necessary either to download the Moa software from github, or, to make use of the integrated version control.
- A number of support scripts & templates depend on Biopython. Consider installing it before starting to use Moa.
- *Python-dev*: the Python development package. A few prerequisites installed by easy\_install try to compile C libraries, and need this. Although all of them have backup, python only, alteratives; from a performace perspective it is probably smart to have this installed:

2.3. Installation 7

```
sudo apt-get install python-dev
```

• *python-yaml*:On ubuntu, this installs a fast YAML parser - using easy\_install or pip might install a slower, python only, version:

```
sudo apt-get install python-yaml
```

## 2.3.2 Python prerequisites

These preregs can be installed manually or with easy\_install or pip:

- pyyaml (unless already installed)
- Jinja2
- · Ruffus
- gitpython
- 'unittest2 http://pypi.python.org/pypi/unittest2'\_
- 'lockfile http://pypi.python.org/pypi/lockfile'\_

Not part of the list of prerequisites are the following libraries, which you'll only need if you are planning to run the web interface:

- ElementTree
- Markdown

#### 2.3.3 Bioinformatics tools

Each of the wrapped tools requires the tools to be present. Usually, Moa expects all tools to be present & executable on the system PATH. The standard Moa distribution comes with wrappers for:

- Blast
- BWA
- Bowtie
- Soap

and many more

## 2.3.4 Installing git (from github)

Moa is hosted on and can be installed from github:

```
cd ~
git clone git://github.com/mfiers/Moa.git moa
```

## 2.3.5 Configuration

Configuration of Moa is simple, and can be done by sourcing the *moainit* script:

```
. ~/moa/bin/moainit
```

(Note the dot!, alternatively use: source ~/moa/bin/moainit)

It is probably a good idea to add this line to your ~/.bashrc for future sessions.

Moa should now work, try moa –help.

If your default python version is NOT *python2.6* or *python2.7* there are a few options that you can pursue:

- change the hashbang of the moa script
- define an alias in your ~/.bashrc: alias moa='python2.6 moa'
- create a symlink to python2.6 in your ~/bin directory and make sure that that is first in your path.

## 2.3.6 Installing the web interface

<Directory /opt/moa/www/html>
 Order allow,deny
 Allow from all
 Options +Indexes

</Directory>

Note - this is experimental - you will probably need to fiddle a little with the configuration to get it working. Start with installing apache2.

Then - assuming that: \* Your Moa work directory is under /home/moa/work \* Your Moa is installed in /opt/moa Create a file in /etc/apache2/conf.d/moa.conf with the following approximate contents:

```
Alias /moa/data /home/moa/work
<Directory /home/moa/work>
   Options +Indexes +FollowSymLinks
   Order allow, deny
  Allow from all
   SetEnv MOADATAROOT /home/moa/work
   SetEnv MOAWEBROOT /moa/data
   IndexOptions FoldersFirst SuppressRules HTMLTable IconHeight=24 SuppressHTMLPreamble
  HeaderName /moa/cgi/indexHeader.cgi
   ReadmeName /moa/html/indexFooter.html
</Directory>
ScriptAlias /moa/cgi/ /opt/moa/www/cgi/
<Directory /opt/moa/www/cgi/>
    AddType text/html .cgi
   Order allow, deny
   Allow from all
    SetEnv MOABASE /opt/moa
</Directory>
Alias /moa/html/ /opt/moa/www/html/
```

2.3. Installation 9

You might want to check the #! of /opt/moa/www/cgi/indexHeader.cgi depending on your system configuration. Restart apache and it should work

## 2.4 Three core templates

Moa comes with a list of templates (see *templates*). The three most important, flexible templates of these that allow you to embed custom code (called *process*) in your project are:

simple:

Simply executes process as a bash one-liner

map:

Takes a set of in- and output files and executes the custom commands for each in- and output file (using the Jinja2 template language).

reduce:

Takes a set of input files and a single output file and executes the custom commands with all input file, generating the output files.

Since *simple*, *map* and *reduce* have proven to be quite central to how Moa operates they come with their own shortcut commands (*moa simple*, *moa map* and *moa reduce*). These command query the user directly for the parameters instead of having to define this manually.

For example, a simple job:

```
$ mkdir simple_test && cd simple_test
$ moa simple -t 'Generate some files'
process:
> for x in 'seq 1 5'; do touch test.$x; done
$ moa run
$ ls
test.1 test.2 test.3 test.4 test.5
```

Note that you can make your *process* as complicated as you like. Alternatively, you can write a script that you call from *process*.

A map job would work like this:

```
$ mkdir ../map_test && cd ../map_test
$ moa map -t 'Map some files'
process:
> echo {{ input }} ; echo {{ input }} > {{ output }}
input:
> ../simple_test/test.*
output:
> ./out.*
$ moa run
../simple test/test.3
../simple_test/test.1
../simple_test/test.5
../simple_test/test.2
../simple_test/test.
Moa: Success executing "run" (<1 sec)
$ ls
out.1 out.2 out.3 out.4 out.5
```

```
$ cat out.1
../simple_test/test.1
```

Moa tracks which input file generates which outputfile. So, if you would like to repeat one of the jobs - you'll need to delete the output file & rerun *moa*:

```
$ rm out.3
$ moa run
../simple_test/test.3
Moa: Success executing "run" (<1 sec)
And a reduce example:
$ mkdir ../reduce test && cd ../reduce test
$ moa reduce -t 'Reduce some files'
process:
> echo {{ " ".join(input) }} >> {{ output }}
input:
> ../map_test/out.*
output:
> ./reduce_out
$ moa run
Moa: Success executing "run" (<1 sec)
$ ls
reduce_out
$ cat reduce_out
../map_test/out.1 ../map_test/out.3 ../map_test/out.4 ../map_test/out.5 ../map_test/out.
```

## 2.5 Synchronizing jobs

It is quite often usefull to repeat a jobs on a number of different input files. For simple operations, one liners, this ca be accomplished using *moa map*. More complex operations, or those requiring a template other than *map* can be replicated using job syncronization. Assume you have a set of fastq libraries, each in it's own directory:

```
./fq/set1/set1_1.fq
./fq/set1/set1_2.fq
./fq/set2/set2_1.fq
./fq/set2/set2_2.fq
./fq/set3/set3_1.fq
./fq/set3/set3_2.fq
```

And you want to run a bowtie alignment for each separately. The approach to take is to create a directory containing all alignments:

```
mkdir bowtie cd bowtie
```

and, in that directory, create one job running bowtie, in a directory named **exactly** as the input directories:

```
mkdir set1
cd set1
moa new bowtie -t 'run bowtie for {{_}}'
```

Note the magic variable {{\_}}}. This variable is replaced by the name of the current directory. So when running *moa show*, the title would show up as "run bowtie for set1". This magic variable can be used

in all variables, and we'll use it here to set this job up in such a way that it can be reused for the other datasets:

```
set moa fq_forward_input='../../fq/{{_}}/*_1.fq'
# .. configure the remaining variables
```

Now - we replicate this directory in the following manner. We'll move one directory up, to the *bowtie* directory, and create a *sync* job:

```
cd ..
moa new sync -t 'run bowtie for all fq datasets'
moa set source=../fq/
```

The sync template keeps directories synchronized, based on the *source* directory. If you now run *moa run* in the *bowtie* directory, two more directories will be created: *set2* and *set3*, each containing a verbatim copy of the original bowtie job created.

If, at a certain moment you obtain more fastq datasets:

```
./fq/set4/set4_1.fq
./fq/set4/set4_2.fq
```

you can repeat *moa run* in the *./bowtie* sync directory, and a new directory will be created. Note that the *sync* template will not remove directories. Also if you want to update the configuration of the syncronized bowtie jobs, you only need to change the configuration in one directory, run *moa run* again in the *./bowtie* directory and the configuration is synchronized across all jobs.

NOTE: both the software and the manual are under development. Expect things to change.

## 2.6 How to write a template

A MOA template is made up of a .moa file and a .jinja2 (or .mk) file.

The .moa file mainly contains input-output file sets and parameter options used for the bash command(s). Some of these options have default values which the user can change while constructing the job.

The .jinja2 file includes information to structure the command(s). It is written in jinja, which is a templating language for python and is simple to write and easy to understand.

These files are used by the backend, currently *ruffus*, that manages file set and parameter dependencies to make pipelines and render commands to the bash prompt. Initially, *GNU make* was the backend used. It is very powerful but some of its limitations and its complexity led to including *ruffus* as an option for the backend as well.

The easiest way to write a moa template is to edit an existing template to suit your requirements. This involves understanding the parts of an existing template.

The bwa\_aln template is used as an example below. Just as a background, the *bwa aln* command takes a FASTQ file as input and aligns it to a reference genome that was previously indexed. The output is a sai file with the alignments.

The bwa\_aln.moa file has some main components:

Backend

```
backend: ruff
```

This is 'ruff' which means that ruffus is used in the python script at a lower level to read the template .moa and .jinja2 file, and render the corresponding commands to the bash prompt.

#### Commands

```
commands:
    run:
    mode: map
    help: run bwa aln
    clean:
    mode: simple
    help: Remove all job data, not the Moa job itself, note that this must be imple
```

This indicates the function names that you will later define. In the example above, there are 2 commands- run and clean, so moa run or moa clean on the command prompt in the job directory would execute these functions.

#### • Filesets

```
filesets:
   input:
     category: input
     extension: fq
   help: Fastq input files
     glob: '*'
     optional: false
     type: set
   output:
     category: output
     dir: .
     extension: sai
     glob: '{{ input_glob }}'
     source: input
     type: map
```

Like the name, each filesets refer to a set of files in a single directory. The bwa\_aln template shows 2 filesets: input and output.

- Category: is essentially used to separate input from output.
- Extension: refers to the type of file(s) required or generated.
- Glob: searches for files with a specified pattern. Moa, by default (glob= \*) automatically processes all files of the specified input extension in the directory specified. By specifying a glob, Moa will only process those files whose name pattern matches what is in the glob.
- Type: refers to the data type of the fileset or parameter.

A fileset can either be of set or map type. The type set refers to a simple set of files in a directory. The type map refers to a set of files that are linked to what their source value is. In the above code, the output fileset is mapped to the input fileset.

- *Dir*: the directory of the output fileset is '.', which means that the output files will be placed in the current working directory.
- Parameter category order

#### Parameters

```
mismatch_penalty:
  category: ''
  default: 3
  help: mismatch penalty
  optional: true
  type: integer
```

They are the variables/options that specify a command.

- Category:
- Default: is the value that is used by default if not changed by the user.
- *Optional*: specifies if it is necessary for the user to fill in a value for the variable. If optional is false, the user has to indicate a value for the parameter in order to execute the job.
- Type: specifies the data type of the variable eg. integer, string, boolean.
- Moa id

```
moa id: bwa aln
```

is supposed to be the same as the filename. Ideally something descriptive (eg. bwa\_aln). This is used to later link to the other template file.

The other template file is "bwa\_aln.jinja2" which is written in jinja, a templating language for python. *Note that the jinja2 file name is the same as the moa file name.* 

Important features of the bwa\_aln.jinja2 file are:

• The three hash's (###) specify the start of a function and are followed by the function name. In our bwa\_aln example, we have defined 2 funtions: run and clean.

```
### run
```

• This defination is followed by a set of commands which you would want to be executed when you type moa run or moa\_clean in the bwa\_aln job directory. The commands in our example file look the same as what you would put in the command prompt but the values of the parameters are bought from the .moa file and hence it's value is replaced by the parameter name.

```
bwa aln {{db}}
    -n {{edit_dist_missing_prob}}
    .
    .
    .
    .
    {{ input }}
    -f {{ output}}
```

• It is also possible to add if-else statements or other computing blocks in accordance with the design language.

```
{% if color_space %} -c {% endif %}
```

## 2.7 Command reference

#### 2.7.1 moa!

Create a 'simple' job from the last command issued. Set the *process* parameter to the last issued command. If a moa job exists in the current directory, then the *process* parameter is set without questions. (even if the Moa job in question does not use the *process* parameter). If no moa job exists, a *simple* job is created first. *Note:* This works only when using *bash* and if *moainit* is sourced properly. *moainit* defines a bash function *\_moa\_prompt* that is called every time a command is issued (using \$PROMPT\_COMMAND). The *\_moa\_prompt* function takes the last command from the bash history and stores it in ~/.config/moa/last.command. Additionally, the *\_moa\_prompt* function stores all commands issued in a Moa directory in .moa/local\_bash\_history.

#### optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

--profile Run the profiler-f, --force Force this action

**-t TITLE, --title TITLE** A title for this job

#### 2.7.2 moa archive

Archive a job, or tree with jobs for later reuse. This command stores only those files that are necessary for execution of this job, that is: templates & configuration. In & output files, and any other file are ignored. An exception to this are all files that start with 'moa. If the *name* is omitted, it is derived from the *jobid* parameter. It is possible to run this command recursively with the *-r* parameter - in which case all (moa job containing) subdirectories are included in the archive.

positional arguments: name archive name

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseShow debugging output

--profile Run the profiler-f, --force Force this action

-s, --sync Alternative approach to deal with sync type jobs - only include

\_ref directories

**-t, --template** Store this archive as a template

## 2.7.3 moa blog

Add an entry to the blog job (Blog.md) Allows a user to maintain a blog for this job (in Blog.md). Use as follows:: \$ moa blog Enter your blog message (ctrl-d on an empty line to finish) ... enter your message here .. [ctrl-d] Note: the ctrl-d needs to be given on an empty line. The text is appended to moa.desciption. In the web interface this is converted to Markdown\_. .. \_Markdown: http://daringfireball.net/projects/markdown/ markdown.

#### optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

## 2.7.4 moa change

Add entry to CHANGELOG.md This function allows the user to add an entry to CHANGELOG.md (including a timestamp). Use it as follows:: \$ moa change Enter your changelog message (ctrl-d on an empty line to finish) ... enter your message here .. [ctrl-d] Note: the ctrl-d needs to be given on an empty line. The text is appended to moa.desciption. In the web interface this is converted to Markdown\_. .. \_Markdown: http://daringfireball.net/projects/markdown/ markdown. Note the same can be achieved by specifying the -m parameter (before the command - for example: moa -m 'intelligent remark' set ...

#### optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

#### 2.7.5 moa cp

Copy a moa job, or a tree with jobs (with -r). moa cp copies only those files defining a job: the template files and the job configuration. Additionally, all files in the moa directory that start with *moa*. (for example *moa.description* are copied as well. Data and log files are not copied!. If used in conjunction with the -r (recursive) flag the complete tree is copied.

positional arguments: from copy from this path to copy to this path

#### optional arguments:

**-h, --help** show this help message and exit

**-r, --recursive** Run this job recursively

-v, --verbose Show debugging output

**--profile** Run the profiler

-o, --overwrite if the target dir exists - overwrite (insteadof copying into that dir

#### 2.7.6 moa err

Show the stderr of the most recently executed moa job

## optional arguments:

**-h, --help** show this help message and exit

**-r, --recursive** Run this job recursively

**-v, --verbose** Show debugging output

**--profile** Run the profiler

## 2.7.7 moa files

Show in and output files for this job Display a list of all files discovered (for input & prerequisite type filesets) and inferred from these for map type filesets.

#### optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseShow debugging output

**--profile** Run the profiler

#### 2.7.8 moa kill

Kill a running job. This command checks if a job is running. If so - it tries to kill it by sending SIGKILL (-9) to the job.

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseRun this job recursivelyShow debugging output

**--profile** Run the profiler

#### 2.7.9 moa list

Lists all known templates Print a list of all templates known to this moa installation. This includes locally installed templates as well.

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseRun this job recursivelyShow debugging output

**--profile** Run the profiler

**-d** Print a short template description

#### 2.7.10 moa lock

Lock a job - prevent execution

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

## 2.7.11 moa log

Show activity log Shows a log of moa commands executed. Only commands with an impact on the pipeline are logged, such as *moa run & moa set*.

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

## 2.7.12 moa map

create an adhoc moa 'map' job Moa will query the user for process, input & output files. An example session

## optional arguments:

**-h, --help** show this help message and exit

**-r, --recursive** Run this job recursively

**-v, --verbose** Show debugging output

--profile Run the profiler-f, --force Force this action

-t TITLE, --title TITLE A title for this job

#### 2.7.13 moa mv

Move, rename or renumber a moa job.

positional arguments: from copy from this path to copy to this path

optional arguments:

**-h, --help** show this help message and exit

**-r, --recursive** Run this job recursively

**-v, --verbose** Show debugging output

**--profile** Run the profiler

#### 2.7.14 moa new

Create a new job. This command creates a new job with the specified template in the current directory. If the directory already contains a job it needs to be forced using '-f'. It is possible to define arguments for the job on the commandline using KEY=VALUE after the template. Note: do not use spaces around the '=' sign. Use quotes if you need spaces in variables (KEY='two values')

**positional arguments:** template name of the template to use for this moa job parameter arguments for this job, specifyas KEY=VALUE without

spaces

#### optional arguments:

**-h, --help** show this help message and exit

**-r, --recursive** Run this job recursively

**-v, --verbose** Show debugging output

--profile Run the profiler-f, --force Force this action

-d DIRECTORY, --directory DIRECTORY directory to create the job in

-t TITLE, --title TITLE mandatory job title

## 2.7.15 moa out

Show the stdout of the most recently executed moa job

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseRun this job recursivelyShow debugging output

**--profile** Run the profiler

## 2.7.16 moa pause

Pause a running job

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

## 2.7.17 moa postcommand

Execute 'postcommand'

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

## 2.7.18 moa precommand

Execute 'precommand'

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

#### 2.7.19 moa readme

Edit the README.md file for this job You could, obviously, also edit the file yourself - this is a mere shortcut to try to stimulate you in maintaining one

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

#### 2.7.20 moa refresh

Refresh the template Reload the template from the original repository.

#### optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseShow debugging output

**--profile** Run the profiler

#### 2.7.21 moa resume

Resume a running job

#### optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

#### 2.7.22 moa set

Set one or more variables This command can be used in two ways. In its first form both parameter key and value are defined on the command line: *moa set KEY=VALUE*. Note that the command line will be processed by bash, which can either create complications or prove very useful. Take care to escape variables that you do not want to be expandend and use single quotes where necessary. For example, to include a space in a variable: *moa set KEY='VALUE WITH SPACES'*. Alternative use of the set command is by just specifying the key: 'moa set PARAMETER\_NAME', in which case Moa will prompt the user enter a value - circumventing problems with bash interpretation.

**positional arguments:** parameter arguments for this job, specifyas KEY=VALUE without spaces **optional arguments:** 

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseShow debugging output

--profile Run the profiler-f, --force Force this action

#### 2.7.23 moa show

Show all parameters know to this job. Parameters in **bold** are specifically configured for this job (as opposed to those parameters that are set to their default value). Parameters in red are not configured, but need to be for the template to operate. Parameters in blue are not configured either, but are optional.

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

**-u** show unrendered values (when using inline parameters)

**-R** show recursively defined parameters not specified by the local

template

**-p** show private parameters

-a show all parameters

## 2.7.24 moa simple

Create a 'simple' adhoc job. Simple meaning that no in or output files are tracked. Moa will query you for a command to execute (the *process* parameter).

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

--profile Run the profiler-f, --force Force this action

-t TITLE, --title TITLE A title for this job

## 2.7.25 moa status

Show job status Print a short status of the job, including configuration

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseShow debugging output

**--profile** Run the profiler

**-u** show unrendered values (when using inline parameters)

**-R** show recursively defined parameters not specified by the local

template

**-p** show private parameters

-a show all parameters

#### 2.7.26 moa test

Test the job parameters

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive Run this job recursively-v, --verbose Show debugging output

**--profile** Run the profiler

#### 2.7.27 moa tree

Show a directory tree and job status

positional arguments: filter show only directories that match this filter

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseRun this job recursivelyShow debugging output

**--profile** Run the profiler

-a, -all

## 2.7.28 moa unlock

Unlock a job - allow execution

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseShow debugging output

**--profile** Run the profiler

## 2.7.29 moa unset

Remove a parameter from the configuration Remove a configured parameter from this job. In the parameter was defined by the job template, it reverts back to the default value. If it was an ad-hoc parameter, it is lost from the configuration.

positional arguments: parameter parameter to unset

optional arguments:

**-h, --help** show this help message and exit

**-r, --recursive** Run this job recursively

**-v, --verbose** Show debugging output

**--profile** Run the profiler

## 2.7.30 moa version

print moa version number

## optional arguments:

**-h, --help** show this help message and exit

-r, --recursive-v, --verboseShow debugging output

**--profile** Run the profiler

## 2.7.31 msp

moa set process

Usage:

msp

this is an alias for the often used:

```
moa set process
```

# 2.8 Templates

Contents:

## 2.8.1 abyss\_pe

Run Abysspe

#### **Commands**

```
clean Remove all job datarun Execute abysspe in paired-end mode
```

#### **Filesets**

```
fq_forward fastq input files directory - forward
fq_reverse fastq input files directory - reverse
```

```
type: map
source: fq_forward
category: input
optional: True
pattern: */*_2.fq
```

output soap denovo output file

```
type: single
category: output
optional: True
pattern: {}
```

#### **Parameters**

joinpairs number of pairs needed to consider joining two contigs

2.8. Templates 25

type: integer default: 10 optional: True

kmer kmer size

type: integer default: 31 optional: True

threads no threads to use

type: integer default: 3 optional: True

## miscellaneous

Backend ruff

**Author** Mark Fiers

**Creation date** Mon, 21 Nov 2011 12:47:16

Modification date Mon, 21 Nov 2011 12:47:22

## 2.8.2 abyss\_se

Run Abysspe

#### **Commands**

clean Remove all job data

run Execute abyss se

#### **Filesets**

input fastq input files directory

output soap denovo output file

type: single category: output optional: True

```
pattern: {}
```

#### **Parameters**

kmer kmer size

type: integer default: 31 optional: True

threads no threads to use

type: integer default: 3 optional: True

#### miscellaneous

**Backend** ruff

**Author** Mark Fiers

**Creation date** Mon, 21 Nov 2011 12:47:16

Modification date Mon, 21 Nov 2011 12:47:22

## 2.8.3 adhoc

## Execute an ad hoc analysis

The adhoc template assists in running one-liners - possibly on a set of input files

#### Commands

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

#### **Filesets**

input Input files for adhoc

2.8. Templates 27

#### **Parameters**

#### mode

operation mode: *seq*, sequential: process the input files one by one; *par*, parallel: process the input files in parallel (use with *-j*); *all*: process all input files at once (use \$^ in adhoc\_process) and *simple*: Ignore input files, just execute adhoc\_process once.

type: set default: simple optional: True

name\_sed A sed expression which can be used to derive the output file name for each input file (excluding the path). The sed expression is executed for each input file name, and the result is available as \$t in the \$(adhoc\_process) statement. Make sure that you use single quotes when specifying this on the command line

type: string default: s/a/a/ optional: True

## output\_dir Output subdirectory

type: directory default: . optional: True

**process** Command to execute for each input file. The path to the input file is available as \$< and the output file as \$t. (it is not mandatory to use both parameters, for example "cat \$< > output" would concatenate all files into one big file

type: string default: echo "needs a sensbile command" optional: True

touch use touch files to track if input files have changed.

type: set default: T optional: True

#### miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

#### 2.8.4 archroot

## Helper script for a root archive

Helper script for the root of an archive template

#### **Commands**

run no help defined

#### **Parameters**

moa\_archive\_parameters space separated list of parameters to set for this template

```
type: string
default: {}
optional: False
```

#### miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Tue, 17 Apr 2012 10:21:31

Modification date Tue, 17 Apr 2012 10:21:25

#### 2.8.5 bamextract

#### bamextract

Extract a region from a BAM file

#### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.run Extract a region from a BAM file

2.8. Templates 29

#### **Filesets**

## bam BAM input

```
type: single
category: input
optional: False
pattern: {}
```

**regions** List with regions to extract (id seqid start stop)

```
type: single
category: input
optional: True
pattern: {}
```

#### **Parameters**

flank flanking region to extract

```
type: integer
default: 100
optional: {}
```

#### miscellaneous

Backend ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## **2.8.6** bartab

#### **Bartab**

BARTAB - a tool to process sff files

#### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run

## **Parameters**

extra\_parameters extra parameters to feed bartab

```
type: string
default: "
optional: True
```

## **forward\_primer** remove forward primer

```
type: string
default: "
optional: True
```

## in input file for bartab

```
type: file
default: ''
optional: False
```

## map A file mapping barcodes to metadata

```
type: file
default: ''
optional: True
```

## min\_length minimun acceptable sequence length

```
type: integer
default: 50
optional: True
```

## out base output name

```
type: integer
default: bartab
optional: True
```

2.8. Templates 31

qin Quality scores for the input fasta file

```
type: file
default: "
optional: True
```

reverse\_primer remove reverse primer

```
type: string
default: "
optional: True
```

trim Trim barcode

```
type: set
default: T
optional: True
```

#### miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.7 bdbb

#### **Bidirectional best BLAST hit**

Discover the bidirectional best blast hit between two sets of sequences

#### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.run generate a list of bidirectional best blast hits between two databases of sequences

#### **Filesets**

input\_a First multi fasta input set

```
type: single
category: input
optional: False
pattern: */*.fasta
```

# input\_b Second multi fasta input set

```
type: single
category: input
optional: False
pattern: */*.fasta
```

# output List of bidirectional best blasts hits

```
type: map
source: input_a
category: output
optional: True
pattern: */*.list
```

## **Parameters**

eval e value cutoff

type: float default: 1e-10 optional: True

extract Extract the identified sequences from the input fasta files

type: boolean default: False optional: True

nothreads Threads to run blast with with

type: integer default: 4 optional: True

## **protein** Is this a protein set

```
type: boolean
default: False
optional: True
```

**tblastx** If this is a nucleotide set, use tblastx?? (otherwise use blastn)

```
type: boolean
default: F
optional: True
```

#### miscellaneous

```
Backend ruff
Author Mark Fiers
Creation date Wed Nov 10 07:56:48 2010
Modification date unknown
```

# 2.8.8 bfast\_aln

Generate bam format alignments using bfast

## Commands

```
clean Remove all job data, not the Moa job itselfrun run bfast match, localalign, postprocess commands
```

#### **Filesets**

```
fa_input fasta input file
fq_input fastq input files
output_aln

type: map
source: fq_input
category: output
optional: {}
pattern: ./*.aln

output_bam
```

type: map

```
source: fq_input
category: output
optional: {}
pattern: ./*.bam
```

## **Parameters**

algorithm\_colour\_space true -> colour space, false -> NT space

type: boolean default: False optional: True

avg\_mism\_qual Specifies the average mismatch quality

type: integer default: 10 optional: True

extra\_params\_localalign Any extra parameters for the localalign command

type: string default: " optional: True

extra\_params\_match Any extra parameters for the match command

type: string default: " optional: True

extra\_params\_postprocess Any extra parameters for the postprocess command

type: string default: " optional: True

min\_mapping\_qual Specifies to remove low mapping quality alignments

```
type: integer
      default: -2147483648
      optional: True
min_norm_score Specifies to remove low (alignment) scoring alignments
      type: integer
      default: -2147483648
      optional: True
output_format 0 - BAF, 1 - SAM
      type: integer
      default: 1
      optional: True
paired_opp_strands Specifies that paired reads are on opposite strands
      type: boolean
      default: False
      optional: True
pairing_std_dev Specifies the pairing distance standard deviation to examine when recuing
      type: float
      default: 2.0
      optional: True
print_params print program parameters
      type: boolean
      default: False
      optional: True
```

type: integer

thread\_num Specifies the number of threads to use

```
default: 1
optional: True
```

## timing\_information specifies output timing information

```
type: boolean
default: True
optional: True
```

## ungapped\_aln Do ungapped local alignment

```
type: boolean
default: False
optional: True
```

## ungapped\_pairing\_rescue Specifies that ungapped pairing rescue should be performed

```
type: boolean
default: False
optional: True
```

## unpaired\_reads True value specifies that pairing should not be performed

```
type: boolean
default: False
optional: True
```

# usage\_summary Display usage summary (help)

```
type: boolean
default: False
optional: True
```

# which\_strand 0 - consider both strands, 1 - forwards strand only, 2 - reverse strand only

```
type: integer
default: 0
```

```
optional: True
```

## miscellaneous

```
Backend ruff
```

Author Yogini Idnani, Mark Fiers

Creation date Wed Feb 15 10:06:48 2011

Modification date unknown

# 2.8.9 bfast db

Generate db index files for aligning reads with bfast

#### **Commands**

```
clean Remove all job data, not the Moa job itselfrun run bfast fasta2brg and index commands
```

#### **Filesets**

fa\_input fasta input file

## **Parameters**

algorithm\_colour\_space true -> colour space, false -> NT space

type: boolean default: False optional: True

**depth** The depth of the splitting(d). The index will be split into 4<sup>d</sup> parts.

type: integer default: 0 optional: True

extra\_params Any extra parameters

type: string default: ''

```
optional: True
```

**hash\_width** The hash width for the index (recommended from manual = 14)

```
type: integer
default: {}
optional: False
```

index\_num Specifies this is the ith index you are creating

```
type: integer
default: 1
optional: True
```

mask The mask or spaced seed to use.

```
type: string
default: {}
optional: False
```

print\_params print program parameters

```
type: boolean
default: False
optional: True
```

thread\_num Specifies the number of threads to use

```
type: integer
default: 1
optional: True
```

timing\_information specifies output timing information

```
type: boolean
default: True
optional: True
```

usage\_summary Display usage summary (help)

type: boolean default: False optional: True

#### miscellaneous

Backend ruff

Author Yogini Idnani, Mark Fiers

Creation date Wed Feb 15 10:06:48 2011

Modification date unknown

#### 2.8.10 blast

## **Basic Local Alignment Tool**

Wraps BLAST [[Alt90]], probably the most popular similarity search tool in bioinformatics.

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template.

report Generate a text BLAST report.

**run** Running BLAST takes an input directory, determines what sequences are present and executes BLAST on each of these. Moa BLAST is configured to create XML output (as opposed to the standard text based output) in the out directory. The output XML is subsequently converted to GFF3 by the custom blast2gff script (using BioPython). Additionally, a simple text report is created.

## **Filesets**

db Blast database

type: single

category: prerequisite

optional: False pattern: \*/\*

input Directory with the input files for BLAST, in Fasta format

outgff GFF output files

```
type: map
source: input
category: output
optional: True
pattern: gff/*.gff
```

## output XML blast output files

```
type: map
source: input
category: output
optional: True
pattern: out/*.out
```

## **Parameters**

eval e value cutoff

```
type: float
default: 1e-10
optional: True
```

 $\mathbf{gff\_blasthit}$   $(T,^*F^*)$  - export an extra blasthit feature to the created gff, grouping all hsp (match) features.

```
type: set
default: F
optional: True
```

gff\_source source field to use in the gff

```
type: string
default: BLAST
optional: True
```

nohits number of hits to report

```
type: integer
default: 50
```

```
optional: True
```

nothreads threads to run blast with (note the overlap with the Make -j parameter)

```
type: integer
default: 2
optional: True
```

program blast program to use (default: blastn)

```
type: set
default: blastn
optional: True
```

#### miscellaneous

```
Backend ruff
```

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.11 blastdb

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** Takes either a set of fasta files or a single multi-fasta input file and creates a BLAST database.

#### **Filesets**

#### dbname

```
type: map
source: input
category: output
optional: {}
pattern: ./db
```

**input** The file with all input FASTA sequences for the blastdb.

```
type: single
```

```
category: input
optional: False
pattern: */*.fasta
```

#### **Parameters**

```
protein Protein database? (T)rue) or not (F)alse (default: F)
```

```
type: set
default: F
optional: True
```

## miscellaneous

Backend ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

**Modification date** Tue, 03 Jan 2012 15:00:23

## 2.8.12 blat

## Blat

Run BLAT on an set of input files (query) vs a database.

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

## **Parameters**

**db** type of the database (dna, prot or dnax)

```
type: set
default: ''
optional: False
```

db\_id\_list a sorted list of db ids and descriptions, enhances the report generated

```
type: file
```

```
default: "
      optional: True
db_type type of the database (dna, prot or dnax)
      type: set
      default: dna
      optional: True
eval evalue cutoff to select the reported hits on (defaults to 1e-15)
      type: float
      default: 1e-10
      optional: True
gff_source Source field for the generated GFF files
      type: string
      default: "
      optional: False
input_dir source field in the generated gff
      type: directory
      default: "
      optional: False
input_extension extension of the input files
      type: string
      default: fasta
      optional: True
input_file input query file. If this variable is not defined, the combination of blat_input_dir and
      blat_input_extension is used to find a list of input files
      type: file
```

```
default: "
optional: False
```

query\_type type of the query (dna, rna, prot, dnax or rnax)

type: set default: dna optional: True

## miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.13 bowtie

## **Bowtie**

Run BOWTIE on an set of input files (query) vs a database index.

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template **run** *no help defined* 

#### **Filesets**

input Fasta/fastq input files for bowtieoutput Output files

type: map source: input category: output optional: {} pattern: ./\*.bam

## **Parameters**

**db** The (basename of the) bowtie database to use.

```
type: string
default: {}
optional: False
```

extra\_params extra parameters to feed bowtie

```
type: string
default: "
optional: True
```

input\_format Format of the input files

```
type: set
default: fastq
optional: True
```

## miscellaneous

**Backend** ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.14 bowtie\_pe

Run BOWTIE on an set of input files (query) vs a database index.

## **Commands**

clean Remove all job data, not the Moa job itself

finish finish up

**report** Create a report on the results

run Execute soapdenovo in paired-end mode

## **Filesets**

**db** The (basename of the) bowtie database to use.

```
type: single
category: prerequisite
optional: False
pattern: ../20.bowtiedb/db
```

fq\_forward\_input Fastq input files - forward

fq\_reverse\_input Fastq input files - reverse

```
type: map
source: fq_forward_input
category: input
optional: True
pattern: */*_2.fq
```

# output Bam output file

```
type: map

source: fq_forward_input

category: output

optional: {}

pattern: ./*.bam
```

## **Parameters**

extra\_params extra parameters to feed to bowtie

```
type: string
default: "
optional: True
```

input\_format Format of the input files

```
type: set
default: fastq
optional: True
```

lots\_of\_data Keep unmapped reads, unsorted BAM - takes up a lot of space!

type: boolean default: False optional: True

max insertsize Maximum allowed insertsize

type: integer default: 250 optional: True

min\_insertsize Minimum allowed insertsize

type: integer default: 1 optional: True

orientation orientation of the reads, allowed values are fr, rf, ff

type: {}
default: fr
optional: True

## miscellaneous

**Backend** ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.15 bowtie\_se

Run BOWTIE on an set of input files (query) vs a database index.

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template **run** *no help defined* 

## **Filesets**

```
fq_input fastq input files directory
output Bam output file

    type: map
    source: fq_input
    category: output
    optional: {}
```

## **Parameters**

**ebwt\_base** The (basename of the) bowtie database to use.

```
type: string
default: {}
optional: False
```

pattern: ./\*.bam

extra\_params extra parameters to feed to bowtie

```
type: string
default: "
optional: True
```

input\_format Format of the input files

```
type: set
default: fastq
optional: True
```

output\_format Format of the output file

```
type: set
default: bam
optional: True
```

## miscellaneous

Backend ruff

Author Yogini Idnani, Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.16 bowtiedb

## **Bowtie index builder**

Builds a bowtie index from a reference sequence

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run Create the bowtie database

#### **Filesets**

input Input fasta file for the bowtie databaseoutput database name to create

```
type: single
category: output
optional: {}
pattern: db
```

#### **Parameters**

extra\_params any option parameters

```
type: string
default: "
optional: True
```

## miscellaneous

Backend ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

**Modification date** Wed Dec 09 07:56:48 2010

# 2.8.17 bwa\_aln

Use BWA to align a set of fastq reads against a db

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** run bwa aln

# **Filesets**

```
input Fastq input files
output

    type: map
    source: input
    category: output
    optional: {}
```

pattern: ./\*.sai

## **Parameters**

best\_hits\_stop stop searching when there are >INT equally best hits

```
type: integer
default: {}
optional: True
```

color\_space input sequences are in the color space

```
type: boolean
default: False
optional: True
```

db bwa database to align against

```
type: string
default: {}
optional: False
```

# edit\_dist\_missing\_prob max

type: float
default: {}
optional: True

# gap\_ext\_max

```
type: integer
default: {}
optional: True
```

gap\_ext\_penalty gap extension penalty

```
type: integer
default: {}
optional: True
```

gap\_open\_penalty gap open penalty

```
type: integer
default: {}
optional: True
```

gap\_opens\_max maximum number or fraction of gap opens

```
type: integer
default: {}
optional: True
```

log\_gap\_penalty\_del log-scaled gap penalty for long deletions

```
type: boolean
default: {}
optional: True
```

max\_ext\_long\_del maximum occurrences for extending a long deletion

```
type: integer
```

```
default: {}
      optional: True
max_queue_entry maximum entries in the queue
      type: integer
      default: {}
      optional: True
mismatch_penalty mismatch penalty
      type: integer
      default: {}
      optional: True
no_indel_from_ends do not put an indel within INT bp towards the ends
      type: integer
      default: {}
      optional: True
non_iterative non-iterative mode search for all n-difference hits (slow)
      type: boolean
      default: False
      optional: True
quality_step quality threshold for read trimming down to 35bp
      type: integer
      default: {}
      optional: True
seed_len Seed length
      type: integer
```

default: {}

```
optional: True
```

# seed\_max\_diff Maximum differences in the seed

```
type: integer
default: {}
optional: True
```

# thread\_num number of threads

```
type: integer
default: {}
optional: True
```

#### miscellaneous

Backend ruff

Author Mark Fiers, Yogini Idnani

**Creation date** Wed Nov 10 07:56:48 2010

Modification date unknown

# 2.8.18 bwa\_index

## Bwa index builder

Builds a bwa index from a reference sequence

#### **Commands**

```
clean Remove all job datarun Create the index
```

## **Filesets**

input Input fasta file for the bowtie database

```
type: single
category: input
optional: False
pattern: */*.fasta
```

## **Parameters**

algorithm Algorithm for constructing BWT index. Available options are 'is' and 'bwtsw'

type: string default: is optional: True

**color\_space** input sequences are in the color space

type: boolean default: False optional: True

prefix Name of the bwa index to create

type: string default: db optional: True

## miscellaneous

**Backend** ruff

Author Mark Fiers, Yogini Idnani

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.19 bwa\_sampe

Generate alignments in SAM format given paired end reads

## **Commands**

clean Remove all job data, not the Moa job itselfrun run bwa sampe

## **Filesets**

fq\_forward\_input fastq input files directory - forward
fq\_reverse\_input fastq input files directory - reverse

```
type: map
      source: fq_forward_input
      category: input
      optional: True
      pattern: */*_2.fq
output_bam
      type: map
      source: fq_forward_input
      category: output
      optional: {}
      pattern: ./*.bam
sai_forward_input sai input files - forward
      type: map
      source: fq_forward_input
      category: input
      optional: False
      pattern: */*_1.sai
sai_reverse_input sai input files - reverse files
      type: map
      source: sai_forward_input
      category: input
      optional: True
      pattern: */*_2.sai
Parameters
db bwa database to align against
      type: string
      default: {}
      optional: False
disable_insert_size disable insert size estimate (force -s)
      type: boolean
```

```
default: False optional: True
```

## disable\_SW disable Smith-Waterman for the unmapped mate

```
type: boolean
default: False
optional: True
```

# max\_aln\_out maximum hits to output for paired reads

```
type: integer
default: 3
optional: True
```

## max\_insert\_size maximum insert size

```
type: integer
default: 500
optional: True
```

## max\_occ\_read maximum occurrences for one end

```
type: integer
default: {}
optional: True
```

# max\_out\_discordant\_pairs maximum hits to output for discordant pairs

```
type: integer
default: {}
optional: True
```

# preload\_index preload index into memory (for base-space reads only)

type: boolean default: False

```
optional: True
```

prior\_chimeric\_rate prior of chimeric rate (lower bound)

```
type: integer
default: {}
optional: True
```

#### miscellaneous

```
Backend ruff
```

Author Yogini Idnani, Mark Fiers

**Creation date** Wed Nov 25 17:06:48 2010

Modification date unknown

# 2.8.20 bwa\_samse

Generate alignments in SAM format given single end reads, using both 'bwa samse'.

## **Commands**

```
clean Remove all job data, not the Moa job itself run run bwa samse
```

## **Filesets**

```
fq_input fastq input file
output_bam output bam file
```

```
type: map
source: fq_input
category: output
optional: {}
pattern: ./*.bam
```

sai\_input sai input directory - filenames must correspond to the fastq input files

```
type: map
source: fq_input
category: input
```

optional: False pattern: \*/\*.sai

## **Parameters**

db bwa database to align against

type: string default: '' optional: False

max\_aln\_out Maximum number of alignments to output in the XA tag for reads paired properly

type: integer default: 3 optional: True

## miscellaneous

Backend ruff

Author Yogini Idnani, Mark Fiers

**Creation date** Wed Nov 25 17:06:48 2010

Modification date unknown

## 2.8.21 cdsmatrix

## CdsMatrix

Predicts (prokaryotic) using glimmer3.

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** Generate a matrix of CDS's

#### **Filesets**

input Directory with the cds files for Glimmer3output Output blast files

type: map

```
source: input
category: output
optional: True
pattern: ./*.out
```

reference multi fasta file

```
type: single
category: prerequisite
optional: {}
pattern: */*.fasta
```

table table files

```
type: map
source: input
category: output
optional: True
pattern: ./*.tab
```

## **Parameters**

cutoff score cutoff value - disregards hits below this score

```
type: {}
default: 100
optional: True
```

# miscellaneous

```
Backend ruff
```

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Thu, 21 Jul 2011 20:31:10 +1200

# 2.8.22 cleanFasta

## clean Fasta

Convert files to unix format and convert all characters that are not an A,C,G,T or N to N.

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** Cleanup of a FASTA file (in place!)

#### **Parameters**

cf\_input\_dir Directory with the sequences to run cleanfasta on

```
type: directory
default: ''
optional: False
```

# cf\_input\_extension input file extension

```
type: string
default: fasta
optional: True
```

## $sed\_command$

```
type: string
default: /^>/!s/[^ACGTNacgtn]/N/g
optional: True
```

## miscellaneous

```
Backend gnumakeAuthor Mark FiersCreation date Wed Nov 10 07:56:48 2010Modification date Wed Nov 10 07:56:48 2010
```

# 2.8.23 clustalgroup

#### clustalw

Run clustalw on two sets of sequences

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** run clustalw

## **Parameters**

cwg\_input\_dir This set of sequences to run clustalw on

```
type: directory
default: "
optional: False
```

cwg\_input\_extension Input file extension

```
type: string
default: fasta
optional: True
```

## miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.24 clustalpair

## clustalw

Run clustalw on two sets of sequences

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run run clustalw

## **Parameters**

```
type: directory
default: "
optional: False
```

## input\_dir\_b The set to compare against

```
type: directory
default: "
optional: False
```

## input\_extension Extension of the input files

```
type: string
default: fasta
optional: True
```

#### miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.25 clustalw

#### clustalw

Run clustalw on two sets of sequences

# Commands

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run run clustalw

## **Parameters**

input\_dir\_a This set is compared to the sequences in input\_dir\_b.

```
type: directory
default: "
optional: False
```

```
type: directory
default: "
optional: False
```

# input\_extension Extension of the input files

```
type: string
default: fasta
optional: True
```

## miscellaneous

**Backend** gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.26 concatenate

## Concatenate

Concatenate a set of fasta files into one.

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run no help defined

## **Parameters**

input\_dir Directory with the input data

```
type: directory
default: "
optional: False
```

# input\_extension Extension of the input files

type: string default: fasta optional: True name name of the file, the outputfile will become ./name.fasta

```
type: string
default: "
optional: False
```

## miscellaneous

```
Backend gnumake

Author Mark Fiers

Creation date Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010
```

# 2.8.27 dottup

## **EMBOSS Dottup**

Use dottup (from EMBOSS) to compare two sets of sequences

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** Run dottup

## **Parameters**

input\_dir\_a This set is compared to the sequences in input\_dir\_b.

```
type: directory
default: ''
optional: False
```

input\_dir\_b The set to compare against

```
type: directory
default: "
optional: True
```

input\_extension Extension of the dottup input files

type: string default: fasta optional: True

wordsize Wordsize used to discover similarities between sequences

type: integer default: 8 optional: True

## miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.28 empty

empty

Do nothing...

## **Commands**

## **Parameters**

#### miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Mon Apr 04 16:02:58 2011

Modification date Mon Apr 04 16:03:18 2011

# 2.8.29 fasta2gff

#### **GFF from FASTA**

Derive GFF from a FASTA file, usually to accompany the Sequence for upload to a generic genome browser database.

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

## **Parameters**

f2g\_gffsource Source to be used in the gff

```
type: string
default: ''
optional: False
```

f2g\_input\_dir Directory with the input fasta files

```
type: directory
default: ''
optional: False
```

**f2g\_input\_extension** glob pattern of the fasta files (default: \*.fasta)

```
type: string
default: fasta
optional: True
```

**f2g\_options** options to be passed to the fasta2gff script

```
type: string
default: "
optional: True
```

f2g\_output\_dir Directory with the output gff

```
type: directory
default: ./gff
optional: True
```

## miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.30 fastainfo

## gather information on a set of fasta files

gather info on a set of input files

## Commands

finish create a report

run generate info on each of the input sequences

## **Filesets**

```
input "fastainfo" input files
```

output "fastainfo" raw output files

type: map source: input category: output optional: True pattern: stats/\*.out

stats "fastainfo" collect stat files

type: map source: input category: output optional: True pattern: stats/\*.stat

## **Parameters**

## miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Mon, 11 Jul 2011 15:15:20

Modification date Mon, 11 Jul 2011 15:15:12

# 2.8.31 fastqc

## Run FastQC for fastq QC

Run FastQC on a set a fastq files - quality assessment

### **Commands**

```
finish Run Fastqc
```

finish delegates execution to: report

report Generate a simple fastqc report

run no help defined

#### **Filesets**

input fastqc input files'

**touch** touch files - track if a file has been processed - do not touch this unless you know what you're doing.

type: map source: input category: output optional: True pattern: ./\*.touch

### **Parameters**

output\_dir output directory for the fastQC report

type: dir default: . optional: True

# miscellaneous

Backend ruff

**Author** Mark Fiers

```
Creation date Thu, 28 Apr 2011 09:27:17 +1200 
Modification date Thu, 28 Apr 2011 14:19:04 +1200
```

# 2.8.32 fastx\_clipper

```
run fastx_clipper
```

#### **Commands**

```
clean Remove all job data, not the Moa job itselfrun run fastx_clipper
```

#### **Filesets**

```
input fastq input files directory
output
```

```
type: map
source: input
category: output
optional: {}
pattern: ./*.fq
```

#### **Parameters**

adaptor ADAPTER string. default is CCTTAAGG (dummy adapter).

```
type: string
default: CCTTAAGG
optional: True
```

adaptor\_and\_bases Keep the adapter and N bases after it.

```
type: integer
default: 0
optional: True
```

compress\_output Compress output with GZIP.

```
type: boolean
default: False
```

```
optional: True
```

# debug\_output DEBUG output.

```
type: boolean
default: False
optional: True
```

## help help screen

type: boolean default: False optional: True

**keep\_unknown\_nuc\_seq** keep sequences with unknown (N) nucleotides. default is to discard such sequences.

type: boolean default: False optional: True

out\_adaptor\_only\_seq Report Adapter-Only sequences.

type: boolean default: False optional: True

rm\_clipped\_seq Discard clipped sequences (i.e. - keep only sequences which did not contained the adapter).

type: boolean default: False optional: True

rm\_non\_clipped\_seq Discard non-clipped sequences (i.e. - keep only sequences which contained the adapter).

type: boolean

default: False optional: True

rm\_short\_seq discard sequences shorter than N nucleotides. default is 5.

type: integer default: 5 optional: True

**verbose** Verbose - report number of sequences. If [-o] is specified, report will be printed to STDOUT. If [-o] is not specified (and output goes to STDOUT), report will be printed to STDERR.

type: boolean default: False optional: True

#### miscellaneous

**Backend** ruff

Author Mark Fiers, Yogini Idnani

Creation date Wed Dec 06 17:06:48 2010

Modification date unknown

# 2.8.33 fastx\_qual\_stats

run fastx\_quality\_stats, fastq\_quality\_boxplot\_graph.sh and fastx\_nucleotide\_distribution\_graph.sh

#### **Commands**

clean Remove all job data, not the Moa job itself

run run fastx\_quality\_stats, fastq\_quality\_boxplot\_graph.sh and fastx\_nucleotide\_distribution\_graph.sh

### **Filesets**

## boxplot\_output

type: map source: input category: output optional: {}

```
pattern: ./*.png
input fastq input files directory
nuc_distr_output
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./*.png
qual_output
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./*.txt
Parameters
gen_postScript_file Generate PostScript (.PS) file. Default is PNG image.
      type: boolean
      default: False
      optional: True
graph_title Title - will be plotted on the graph.
      type: string
      default: {{ input_glob }}
      optional: True
help help screen
      type: boolean
      default: False
      optional: True
new_out_format New output format (with more information per nucleotide/cycle)
      type: boolean
      default: False
```

```
optional: True
```

#### miscellaneous

```
Backend ruff
```

Author Mark Fiers, Yogini Idnani

**Creation date** Wed Dec 03 17:06:48 2010

Modification date unknown

# 2.8.34 filterwgs\_pair

Execute a "map22" ad-hoc analysis - two input files, two output files

Filter raw WGS data

#### **Commands**

run Filter WGS data

## **Filesets**

```
input1 forward input fastqinput2 reverse input fastq
```

type: map source: input1 category: input optional: False pattern: \*/\*

## output1 forward output fastq

type: map source: input1 category: output optional: True pattern: ./\*

## output2 reverse output fastq

type: map

```
source: input1
category: output
optional: True
pattern: ./*
```

#### **Parameters**

adapters Fasta file with the adapter sequences to trim

```
type: file
default: {}
optional: False
```

minlen Minimum remaining sequence length

```
type: int
default: 50
optional: True
```

qual quality threshold causing trimming

```
type: int
default: 13
optional: True
```

### title

```
type: {}
default: Filter paired fastq files using fastq-mcf
optional: {}
```

### miscellaneous

```
Backend ruff
```

**Author** Mark Fiers

**Creation date** Tue Mar 29 16:34:19 2011

**Modification date** Mon, 13 Feb 2012 09:16:36 +1300

# 2.8.35 gather

## gather files

gather a set of files and create hardlinks to. Hardlinks have as advantage that updates are noticed via the timestamp. Hence, make recognizes them.

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** gather files

#### **Parameters**

**g\_input\_dir** list of directories with the input files

```
type: directory
default: "
optional: False
```

g\_input\_pattern glob pattern to download

```
type: string
default: *
optional: True
```

**g\_limit** limit the number of files gathered (with the most recent files first, defaults to 1mln)

```
type: integer
default: 1000000
optional: True
```

g\_name\_sed SED expression to be executed on each file name - allows you to change file names

```
type: string
default: s/a/a/
optional: True
```

**g\_output\_dir** Output subdirectory, defaults to .

```
type: directory
default: .
optional: True
```

**g\_parallel** allow parallel execution (T) or not (**F**). If for example concatenating to one single file, you should not have multiple threads.

```
type: set
default: F
optional: True
```

**g\_powerclean** Do brute force cleaning (T/F). Remove all files, except moa.mk & Makefile when calling make clean. Defaults to F.

```
type: set
default: F
optional: True
```

**g\_process** Command to process the files. If undefined, hardlink the files.

```
type: string
default: In -f $$< $$(g_target)
optional: True</pre>
```

#### miscellaneous

```
Backend gnumake
Author Mark Fiers
Creation date Wed Nov 10 07:56:48 2010
Modification date Wed Nov 10 07:56:48 2010
```

# 2.8.36 genemarks

# ${\bf gene Mark S}$

predict genes using geneMarkS

### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run no help defined

### **Filesets**

input Directory with the input files for Genemarks

### **Parameters**

gff\_source source field to use in the gff. Defaults to "geneMarkS"

type: string

default: genemarkS optional: True

matrix the matrix to use

type: file default: '' optional: True

## miscellaneous

Backend ruff

Author

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.37 getorf

#### Getorf

Predicts open reading frames using the EMBOSS [[emboss]] getorf tool.

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

## **Filesets**

```
gff
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./gff/*.gff
input Input files for getorf
output
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./out/*.out
Parameters
circular Is the sequence linear?
      type: set
      default: N
      optional: True
find What to output? 0: Translation between stop codons, 1: Translation between start & stop codon, 2:
      Nucleotide sequence between stop codons; 3: Nucleotide sequence between start and stop codons.
      Default: 3
      type: set
      default: 3
      optional: True
gff_source source field to use in the gff.
      type: string
```

maxsize maximal nucleotide size of the predicted ORF.

default: getorf optional: True

type: integer default: 1000000 optional: True

minsize minimal nucleotide size of the predicted ORF.

type: integer default: 30 optional: True

table Genetic code to use: 0 Standard; 1 Standard with alternative initiation codons; 2 Vertebrate Mitochondrial; 3 Yeast Mitochondrial; 4 Mold, Protozoan, Coelenterate Mitochondrial and Mycoplasma/Spiroplasma; 5 Invertebrate Mitochondrial; 6 Ciliate Macronuclear and Dasycladacean; 9 Echinoderm Mitochondrial; 10 Euplotid Nuclear; 11 Bacterial; 12 Alternative Yeast Nuclear; 13 Ascidian Mitochondrial; 14 Flatworm Mitochondrial; 15 Blepharisma Macronuclear; 16 Chlorophycean Mitochondrial; 21 Trematode Mitochondrial; 22 Scenedesmus obliquus; 23 Thraustochytrium Mitochondrial.

type: set default: 11 optional: True

#### miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.38 glimmer3

#### Glimmer3

Predicts (prokaryotic) using glimmer3.

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template.

**run** Glimmer3 is a open reading frame discovery program from the EMBOSS [[emboss]] package. It takes a set of input sequences and predicts all open reading frames. Additionally, this template converts the default output (predicted protein sequences) to GFF3.

## **Filesets**

cds CDS output files from glimmer3

```
type: map
source: input
category: output
optional: True
pattern: cds/*.fasta
```

gff GFF output files from glimmer3

```
type: map
source: input
category: output
optional: True
pattern: gff/*.gff
```

**input** Directory with the input files for Glimmer3

output Raw output files from glimmer3

```
type: map
source: input
category: output
optional: True
pattern: out/*.g3
```

pep peptide output files from glimmer3

```
type: map
source: input
category: output
optional: True
pattern: pep/*.fasta
```

### **Parameters**

```
gene_len Minimum gene length (glimmer3 -g/-gene_len)
```

```
type: integer
default: 110
optional: True
```

gff\_source source field to use in the gff. Defaults to "glimmer3"

```
type: string
default: glimmer3
optional: True
```

max\_overlap Maximum overlap, see the glimmer documentation for the -o or -max\_olap parameter

```
type: integer
default: 50
optional: True
```

stop\_codons

```
type: {}
default: tag,tga,taa,nnn,tnn,ann,gnn,cnn
optional: True
```

treshold treshold for calling a gene a gene (glimmer3 -t)

```
type: integer
default: 30
optional: True
```

### miscellaneous

Backend ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.39 gmap

**Gmap** 

Run GMAP on an set of input files (query) vs a database index.

### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

#### **Filesets**

```
align
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./align/*.align
genepred
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./genepred/*.genepred
gff
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./gff/*.gff
gff_invert
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./gff/*.invert.gff
input Sequences to map
raw
      type: map
      source: input
      category: output
      optional: {}
```

pattern: ./raw/\*.raw

### **Parameters**

# **db** Gmap db

```
type: file
default: ''
optional: False
```

# extra\_parameters extra parameters to feed to gmap

```
type: string
default: "
optional: True
```

# gff\_source Source field to use in the output GFF

```
type: string
default: gmap
optional: True
```

# **invert\_gff** Invert the GFF (T/*F*)

```
type: set
default: T
optional: True
```

### miscellaneous

```
Backend ruff
```

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.40 gmapdb

## gmapdb index builder

Builds gmapdb index from a reference sequence

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

### **Filesets**

input The reference sequence to build a gmap database with.

```
type: single
category: input
optional: False
pattern: */*.fasta
```

### **Parameters**

name Name of the gmap index to create

```
type: string
default: gmapdb
optional: True
```

# miscellaneous

```
Backend ruff
```

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.41 gsMapper

### **GSMapper**

Run the Roche GS Reference mapper

## **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

#### **Parameters**

annotation Gene annotation file in the UCSC GenePred format

```
type: file
default: "
optional: True
```

min\_overlap\_ident Minimum identity length in the assembly step

```
type: integer
default: 90
optional: True
```

min\_overlap\_len Minimum overlap length in the assembly step

```
type: integer
default: 40
optional: True
```

name Name identifying this mapping in the output gff

```
type: string
default: "
optional: False
```

reference\_fasta A multifasta file with the reference sequence(s) with the library id.

```
type: file
default: "
optional: True
```

sfffile SFF files with reads to map against the reference sequences

```
type: file
default: "
optional: True
```

### miscellaneous

```
Backend gnumake
```

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.42 h\_blast

### **Hadoop Blast**

Runs BLAST on a hadoop cluster

### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run Similar to a normal blast, but now running on an hadoop cluster

#### **Parameters**

db Location of the blast database

```
type: file
default: ''
optional: False
```

eval e value cutoff

type: float default: 1e-10 optional: True

hadoop\_base location of the hadoop installation

```
type: directory
default: "
optional: False
```

hdfs\_base htfs://SERVER:PORT for the hdfs filesystem, defaults to "hdfs://localhost:9000"

```
type: string
      default: hdfs://localhost:9000
      optional: True
input_dir location of the hadoop installation
      type: directory
      default: "
      optional: False
input_extension input file extension
      type: string
      default: fasta
      optional: True
nohits number of hits to report
      type: integer
      default: 50
      optional: True
nothreads threads to run blast with (note the overlap with the Make -j parameter)
      type: integer
      default: 1
      optional: True
program blast program to use (default: blastn)
```

## miscellaneous

type: set default: blastn optional: True

**Backend** gnumake **Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.43 hagfish

## Run hagfish\_extract & hagfish\_combine

Run the preparatory steps for hagfish

## **Commands**

```
clean remove all Hagfish filesfinish finish up - find gaps - combine plots - create a reportrun Run hagfish
```

### **Filesets**

fasta fasta sequence of the reference

```
type: single
category: prerequisite
optional: False
pattern: {}
```

input "hagfish" input files

output "hagfish" touch files - track what files are done - please do not touch this!

```
type: map
source: input
category: output
optional: True
```

pattern: ./touch/\*.touch

#### **Parameters**

circosbinsize Binsize for generating circos formatted histograms

```
type: int
default: {}
optional: True
```

max\_ok Maximal acceptable insert size for an aligned pair. If omitted, hagfish will make an estimate

```
type: int
default: 0
optional: True
```

min\_ok Minimal acceptable insert size for an aligned pair. If omitted, hagfish will make an estimate

```
type: int
default: 0
optional: True
```

#### miscellaneous

```
Backend ruff
```

**Author** Mark Fiers

**Creation date** Tue Mar 29 16:34:19 2011

**Modification date** Thu, 19 May 2011 20:49:04 +1200

# 2.8.44 kanga

use kanga to align short reads to a reference genome

## Commands

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** run kanga

#### **Filesets**

input\_fasta Fasta input file
output output files

```
type: map
source: rds_input
category: output
optional: True
pattern: ./*.sam
```

output\_bam output files

```
type: map
      source: rds_input
      category: output
      optional: True
      pattern: ./*.bam
output_log output log file
      type: map
      source: rds_input
      category: output
      optional: {}
      pattern: ./*.log.txt
rds_input rds (preprocessed) input files
sfx_input sfx array lookup file
Parameters
color_space process for colorspace (SOLiD)
      type: boolean
      default: False
      optional: True
extra_params any extra parameters
      type: string
      default: "
      optional: True
help print this help and exit
      type: boolean
      default: False
```

optional: True

max\_Ns maximum number of intermediate N's in reads before treating read as unalignable

```
type: integer
      default: 1
      optional: True
max_pair_len accept paired end alignments with apparent length of at most this
      type: integer
      default: 300
      optional: True
min_pair_len accept paired end alignments with apparent length of at least this
      type: integer
      default: 100
      optional: True
no_multireads do not accept multiple reads aligning to the same loci
      type: boolean
      default: False
      optional: True
out_format 0 - CSV loci only, 1 - CSV loci + match sequence, 2 - CSV loci + read sequence, 3 - CSV
      loci + read + match sequence, 4 - UCSC BED, 5 - SAM format
      type: integer
      default: 0
      optional: True
pe_mode 0 - none, 1 - paired ends with recover orphan ends, 2 - paired end no orphan recovery
      type: integer
      default: 0
      optional: True
```

quality fastq quality scoring- 0 - sanger, 1m - Illumina 1.3+, 2 - Solexa < 1.3, 3 - Ignore quality

```
type: integer
default: 3
optional: True
```

thread\_num number of processing threads (0 sets threads to number of CPU cores)

```
type: integer
default: 0
optional: True
```

trim3 trim this number of bases from 3' end of reads when loading raw reads

```
type: integer
default: 0
optional: True
```

trim5 trim this number of bases from 5' end of reads when loading raw reads

```
type: integer
default: 0
optional: True
```

version print version information and exit

```
type: boolean
default: False
optional: True
```

### miscellaneous

```
Backend ruff
```

Author Mark Fiers, Yogini Idnani

**Creation date** Wed Nov 10 07:56:48 2010

Modification date unknown

# 2.8.45 kangar\_pe

use kangar to pre process raw fq reads

### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** run kangar

## **Filesets**

```
fq_forward_input fastq input files - forward - containing the 5' end
fq_reverse_input fastq input files directory - reverse - containing the 3' end
```

```
type: map
source: fq_forward_input
category: input
optional: True
pattern: */*_2.fq
```

## output\_log output log file

```
type: map
source: fq_forward_input
category: output
optional: {}
pattern: ./*.log.txt
```

## rds\_output output rds file

```
type: map
source: fq_forward_input
category: output
optional: True
pattern: ./*.rds
```

#### **Parameters**

extra\_params any extra parameters

```
type: string
default: "
optional: True
```

## **help** print this help and exit

```
type: boolean
default: False
optional: True
```

mode processing mode 0 - single end create, 1 - paired end create, 2 - output statistics 3 - dump as fasta

```
type: integer
default: 0
optional: True
```

quality fastq quality scoring- 0 - sanger, 1m - Illumina 1.3+, 2 - Solexa < 1.3, 3 - Ignore quality

```
type: integer
default: 3
optional: True
```

reads\_num limit number of reads (or dumps) in each input file to this many, 0 if no limit

```
type: integer
default: 0
optional: True
```

rm\_duplicates remove duplicate reads retaining only one

```
type: boolean
default: False
optional: True
```

trim3 trim this number of bases from 3' end of sequence

```
type: integer
default: 0
optional: True
```

trim5 trim this number of bases from 5' end of sequence

```
type: integer
default: 0
optional: True
```

version print version information and exit

```
type: boolean
default: False
optional: True
```

### miscellaneous

```
Backend ruff
```

Author Mark Fiers, Yogini Idnani

**Creation date** Wed Nov 10 07:56:48 2010

Modification date unknown

# 2.8.46 kangar se

use kangar to pre process raw fq single end reads

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** run kangar

## **Filesets**

```
fq_input fastq input files - forward - containing the 5' end
output_log output log file
```

```
type: map
source: fq_input
category: output
optional: {}
pattern: ./*.log.txt
```

rds\_output output rds file

```
type: map
```

```
source: fq_input
category: output
optional: True
pattern: ./*.rds
```

#### **Parameters**

extra\_params any extra parameters

```
type: string
default: ''
optional: True
```

**help** print this help and exit

```
type: boolean
default: False
optional: True
```

mode processing mode 0 - single end create, 1 - paired end create, 2 - output statistics 3 - dump as fasta

```
type: integer
default: 0
optional: True
```

 $\mathbf{quality}$  fastq quality scoring- 0 - sanger, 1m - Illumina 1.3+, 2 - Solexa < 1.3, 3 - Ignore quality

```
type: integer
default: 3
optional: True
```

reads\_num limit number of reads (or dumps) in each input file to this many, 0 if no limit

```
type: integer
default: 0
optional: True
```

rm\_duplicates remove duplicate reads retaining only one

type: boolean default: False optional: True

trim3 trim this number of bases from 3' end of sequence

type: integer default: 0 optional: True

trim5 trim this number of bases from 5' end of sequence

type: integer default: 0 optional: True

version print version information and exit

type: boolean default: False optional: True

## miscellaneous

Backend ruff

Author Mark Fiers, Yogini Idnani

**Creation date** Wed Nov 10 07:56:48 2010

Modification date unknown

## 2.8.47 kangax

use kangax to create the suffix array lookup database for the reference genome

### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run run kangax

## **Filesets**

type: string default: " optional: True

```
input_fasta Fasta input file
output_log output log file
      type: map
      source: input_fasta
      category: output
      optional: {}
      pattern: ./*.log.txt
output_sfx output suffix array lookup
      type: map
      source: input_fasta
      category: output
      optional: {}
      pattern: ./*.sfx
Parameters
block_seq_len generated suffix blocks to hold at most this length (MB) concatenated sequences
      type: integer
      default: 3300
      optional: True
color_space generate for colorspace (SOLiD)
      type: boolean
      default: False
      optional: True
extra_params any extra parameters
```

### help print this help and exit

type: boolean default: False optional: True

## reference\_species reference species

type: string default: '' optional: False

target\_dep generate target file only if missing or older than any independent source files

type: boolean default: False optional: True

version print version information and exit

type: boolean default: False optional: True

### miscellaneous

Backend ruff

Author Mark Fiers, Yogini Idnani

**Creation date** Wed Nov 10 07:56:48 2010

Modification date unknown

# 2.8.48 Iftp

## lftp

Use LFTP to download files. This template has two modi, one is set lftp\_mode to mirror data, in which case both lftp\_url and lftp\_pattern (default \*) are used. The other modus is lftp\_mode=get, when one file defined by lftp\_url is downloaded. In the mirror mode it is possible to download only those files that are newer as the files already downloaded by using the lftp\_timestamp parameter

### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** execute the download

### **Parameters**

dos2unix Run dos2unix to prevent problems with possible dos text files

```
type: set
default: F
optional: True
```

get\_name target name of the file to download

```
type: string
default: "
optional: True
```

**lftp\_output\_dir** subdir to create & write all output to. If not defined, data will be downloaded to directory containing the Makefile

```
type: directory
default: .
optional: True
```

**lock** Lock this job after running. This means that you will have to manually unlock the job before lftp actually reruns. This is a good choice if your downloading large datasets or have a slow connection

```
type: set
default: T
optional: True
```

**mode** Mode of operation - mirror or get. Mirror enables timestamping. Get just gets a single file. If using get, consider setting depend\_lftp\_timestamp to F. When using get, the full url should be in lftp\_url. lftp\_pattern is ignored. Defaults to mirror.

```
type: set
default: get
optional: True
```

**noclean** set of files not to be deleted by the powerclean

```
type: string
default: moa.mk Makefile
optional: True
```

**pass** password for the remote site, note that this can be defined on the commandline using: make lftp\_pass=PASSWORD

```
type: password
default: "
optional: True
```

pattern glob pattern to download

```
type: string
default: '*'
optional: True
```

**powerclean** Do brute force cleaning (T/F). Remove all files, except moa.mk & Makefile when calling make clean. Defaults to F.

```
type: set
default: F
optional: True
```

**timestamp** Depend on lftp to decide if a file needs updating, else a touchfile is created that you need to delete or touch before updating (T/F)

```
type: set
default: F
optional: True
```

url The base url to download from

```
type: string
default: "
optional: True
```

user username for the remote site

```
type: string
default: ''
optional: True
```

### miscellaneous

```
Backend gnumake

Author Mark Fiers

Creation date Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010
```

# 2.8.49 map

## Execute a "map" ad-hoc analysis

Execute one command, on a number of input files.

### Commands

```
run no help defined
```

### **Filesets**

```
input "map" input files

output "map" output files

type: map
    source: input
    category: output
    optional: True
    pattern: ./*
```

### **Parameters**

## dummy do a dummy run

type: boolean default: False optional: True

## **process** The command to execute

```
type: string
default: True
optional: False
```

### miscellaneous

```
Backend ruff
Author Mark Fiers
Creation date Tue Mar 29 16:34:19 2011
Modification date Wed Mar 30 06:02:01 2011
```

# 2.8.50 map2

## Execute a "map2" ad-hoc analysis

Execute one command, on a number of input files.

### Commands

```
run no help defined
```

#### **Filesets**

```
input1 "map" input files set 1
input2 "map" input files set 2

type: map
source: input1
category: input
optional: False
pattern: */*
```

# output "map" output files

```
type: map
source: input1
category: output
optional: True
pattern: ./*
```

### **Parameters**

**process** The command to execute

type: string default: True optional: False

#### miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Tue Mar 29 16:34:19 2011

Modification date Wed Mar 30 06:02:01 2011

# 2.8.51 map22

# Execute a "map22" ad-hoc analysis - two input files, two output files

Execute one command, on a number of input files.

#### **Commands**

run no help defined

#### **Filesets**

```
input1 "map" input files set 1input2 "map" input files set 2
```

type: map source: input1 category: input optional: False pattern: \*/\*

## output1 "map" output files

type: map source: input1 category: output

```
optional: True pattern: ./*
```

### output2 "map" output files

```
type: map
source: input1
category: output
optional: True
pattern: ./*
```

#### **Parameters**

**process** The command to execute

```
type: string
default: True
optional: False
```

#### miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Tue Mar 29 16:34:19 2011

Modification date Wed Mar 30 06:02:01 2011

# 2.8.52 maq\_fasta2bfa

# Convert fasta to bfa

Converts a FASTA file to MAQ format for use with a BFA a maq\_fasta2bfa index from a reference sequence

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

#### **Filesets**

bfa

```
type: map
source: input
category: output
optional: {}
pattern: ./bfa/*.bfa
input input FASTA files
```

#### **Parameters**

#### miscellaneous

```
Backend gnumake
Author Mark Fiers
Creation date Wed Nov 10 07:56:48 2010
Modification date Wed Nov 10 07:56:48 2010
```

# 2.8.53 maq\_fastq2bfq

# **Convert FASTQ to BFQ**

Converts a FASTQ file to MAQ BFQ format.

### Commands

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

#### **Filesets**

### bfq

```
type: map
source: input
category: output
optional: {}
pattern: ./bfq/*.bfq
input input FASTA files
```

### **Parameters**

#### miscellaneous

```
Backend gnumake

Author Mark Fiers
```

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

### 2.8.54 mag match pair

### MAQ paired ends mapper

Map paired ends to a reference sequence using MAQ

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

#### **Parameters**

**forward\_suffix** Suffix of each forward filename - recognize forward files this way. Note this is not a regular extension, no . is assumed between the filename & suffix

type: string default: \_f.bfq optional: True

maxdist max outer distance for a (non RF) readpair. This applies to illumina matepairs - i.e. short inserts

type: integer default: 250 optional: True

read\_dir directory containing the forward reads

type: string default: '' optional: False

reference Reference bfa file to map the reads to

type: string default: " optional: False

### reverse\_suffix suffix of reverse files

```
type: string
default: _r.bfq
optional: True
```

**RF\_maxdist** max outer distance for an RF readpair (corresponds to the -A parameter). This applies to long insert illumina pairs

```
type: integer
default: 15000
optional: True
```

#### miscellaneous

```
Author Mark Fiers

Creation date Wed Nov 10 07:56:48 2010
```

Modification date Wed Nov 10 07:56:48 2010

# 2.8.55 maq\_pe

Generate alignments in SAM format given paired end reads using Maq.

#### **Commands**

```
clean Remove all job data, not the Moa job itselfrun run maq's fasta2bfa, fastq2bfq and map.
```

#### **Filesets**

bam\_output bam alignment output file

```
type: map
source: fq_forward_input
category: output
optional: {}
pattern: ./*.bam
```

**bfa\_output** BFA Index name

```
type: single
category: other
optional: {}
pattern: {}
```

# bfq\_forward\_output bfq files - forward files

```
type: map
source: fq_forward_input
category: output
optional: {}
pattern: ./*_1.bfq
```

# bfq\_reverse\_output bfq files - reverse files

```
type: map
source: fq_forward_input
category: output
optional: {}
pattern: ./*_2.bfq
```

 $fa\_input$  directory with reference fasta file name

 $\label{prop:continuous} \textbf{fq\_forward\_input} \ \ \text{fastq input files directory - forward files}$ 

fq\_reverse\_input fastq input files directory - reverse files

```
type: map

source: fq_forward_input

category: input

optional: {}

pattern: */*_2.fq
```

## map\_output maq map output files

```
type: map
source: fq_forward_input
category: output
optional: {}
pattern: ./*.map
```

### **Parameters**

disable\_sw disable Smith-Waterman alignment

```
type: boolean
default: False
optional: True
```

extra\_parameters Any extra parameters

```
type: string
default: "
optional: True
```

first\_read\_len length of the first read (<=127)s

```
type: integer
default: 0
optional: True
```

match\_in\_colorspace match in the colorspace

```
type: boolean
default: False
optional: True
```

max\_dist\_read\_pairs max distance between two paired reads s

```
type: integer
default: 250
optional: True
```

max\_dist\_RF\_read\_pairs max distance between two RF paired reads s

```
type: integer
default: 0
optional: True
```

max\_mismatch\_qual\_sum maximum allowed sum of qualities of mismatches

```
type: integer
default: 70
optional: True
```

max\_num\_hits\_out max number of hits to output. >512 for all 01 hits.

```
type: integer
default: 250
optional: True
```

num\_mismatch\_24bp number of mismatches in the first 24bp

```
type: integer
default: 2
optional: True
```

read\_ref\_diff\_rate rate of difference between reads and references

```
type: float
default: 0.001
optional: True
```

sec\_read\_len length of the second read (<=127)s</pre>

```
type: integer
default: 0
optional: True
```

trim\_all\_reads trim all reads (usually not recommended)

```
type: boolean
default: False
optional: True
```

### miscellaneous

```
Backend ruff
Author Mark Fiers, Yogini Idnani
Creation date Wed Dec 03 17:06:48 2010
Modification date unknown
```

# 2.8.56 maq\_se

Generate alignments in SAM format given single end reads using Maq.

#### **Commands**

```
clean Remove all job data, not the Moa job itself run run maq's fasta2bfa, fastq2bfq and map.
```

#### **Filesets**

bam\_output bam alignment output file

```
type: map
source: fq_input
category: output
optional: {}
pattern: ./*.bam
```

### **bfa\_output** BFA Index name

```
type: single
category: other
optional: {}
pattern: {}
```

### **bfq\_output** bfq files - forward files

```
type: map
source: fq_input
category: output
optional: {}
pattern: ./*.bfq
```

```
fa_input directory with reference fasta file name
fq_input fastq input files
map_output maq map output files

type: map
source: fo_input
```

source: fq\_input category: output optional: {} pattern: ./\*.map

#### **Parameters**

disable\_sw disable Smith-Waterman alignment

type: boolean default: False optional: True

extra\_parameters other parameters

type: string default: " optional: True

match\_in\_colorspace match in the colorspace

type: boolean default: False optional: True

max\_mismatch\_qual\_sum maximum allowed sum of qualities of mismatches

type: integer default: 70 optional: True

max\_num\_hits\_out number of mismatches in the first 24bp

type: integer default: 250 optional: True

### num\_mismatch\_24bp number of mismatches in the first 24bp

type: integer default: 2 optional: True

### read\_ref\_diff\_rate rate of difference between reads and references

type: float default: 0.001 optional: True

# trim\_all\_reads trim all reads (usually not recommended)

type: boolean default: False optional: True

# miscellaneous

Backend ruff

Author Mark Fiers, Yogini Idnani

**Creation date** Wed Dec 02 17:06:48 2010

Modification date unknown

### 2.8.57 moatest

### Unittest template

Not to be used - is used by unitmoatests

### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.run Do nothing - no need to call this.

### **Parameters**

test\_opt test variable

type: string default: konijntje optional: True

txt test variable

type: string default: " optional: False

### miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.58 mummer

#### mummer

Run mummer between two sequences

#### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run Run mummer

### **Filesets**

input Set 1 input fasta filesreference Set 1 input fasta files

#### **Parameters**

base base name for all generated files

```
type: {}
default: out
optional: True
```

**breaklen** Set the distance an alignment extension will attempt to extend poor scoring regions before giving up (default 200)

```
type: integer
default: 200
optional: True
```

genomecenter genome center - used in the AGP file

```
type: {}
default: pflnz
optional: True
```

gff\_source GFF source field

```
type: {}
default: mumscaff
optional: True
```

linker linker sequence for the merged output sequence

```
type: {}
default: NNNNNCTAGCTAGCATGNNNNNN
optional: True
```

matchmode use all matching fragments (max) or only unique matchers (mum)

```
type: set
default: mum
optional: True
```

mum\_plot\_raw plot an alternative visualization where mummer does not attempt to put the sequences
in the correct order

```
type: boolean
default: False
optional: True
```

organism Organism name - used in the AGP file

```
type: {}
default: "
optional: True
```

taxid Taxonomy id - used in the AGP file

```
type: {}
default: "
optional: True
```

### miscellaneous

Backend ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

### 2.8.59 ncbi

#### **Download data from NCBI**

Download a set of sequences from NCBI based on a query string *ncbi\_query* and database *ncbi\_db*. This tempate will run only **once**, after a successful run it creates a lock file that you need to remove to rerun

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** Download from NCBI

#### **Parameters**

db NCBI database

```
type: string
```

```
default: nuccore optional: True
```

**query** NCBI query (for example txid9397[Organism%3Aexp])

type: string default: '' optional: True

**rename\_sequence** try to rename the sequence - note, this does not work if you are downloading more than one sequence

type: boolean default: False optional: True

**sequence\_name** Name of the file to write the downloaded sequences to. Use 'from\_dir' to have the sequence name extracted from the directory name

type: string default: out optional: True

#### miscellaneous

Backend ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

#### 2.8.60 newbler

# Newbler

Run a simple, out of the box, newbler assembly. As an extra feature, this template automatically creates uniquely named links to the two main output fasta files (454AllContigs.fna, 454LargeContigs.fna). This is convenient for subsequence gather steps. The links are named after the directory.

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template.

```
run no help defined
```

#### **Filesets**

input input SFF files

#### **Parameters**

largecontig\_cutoff min length of a contig in 454LargeContigs.fna

```
type: integer
default: ''
optional: True
```

**library\_name** A library identifier for this assembly. This is used to create an extra fasta file, named using this variable, that contain the generated contigs with their ids prepended with the library id.

```
type: string default: (shell\ echo\ 'basename\ (CURDIR)\ |\ sed\ ''s/[\ ///]//g''\ )' optional: True
```

mid\_configuration Mid configuration file to use

```
type: file
default: ''
optional: True
```

mids mids to use for this assembly

```
type: string
default: ''
optional: True
```

min\_identity Minimal overalp identity used during assembly

```
type: integer
```

default: '' optional: True

### miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.61 newjobtest

### Execute a "simple" ad hoc analysis

Execute one command, No in or output files are tracked by Moa.

### **Commands**

run no help defined

#### **Parameters**

**process** The command to execute

type: string default: True optional: False

#### miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Tue Mar 29 16:34:19 2011

Modification date Wed Mar 30 06:02:01 2011

#### 2.8.62 nstretch

#### **Nstretch**

Run NSTRETCH on an set of input files

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

#### **Parameters**

input\_dir input dir with the fasta files

```
type: directory
default: ''
optional: False
```

input\_extension extension of the input files

```
type: string
default: fasta
optional: True
```

**len** minimal number of Ns before its reported (default 10)

```
type: integer
default: 10
optional: True
```

### miscellaneous

```
Backend gnumakeAuthor Mark FiersCreation date Wed Nov 10 07:56:48 2010Modification date Wed Nov 10 07:56:48 2010
```

### 2.8.63 orthomcl

### Run OrthoMCL

Execute one command, No in or output files are tracked by Moa.

### **Commands**

run no help defined

### **Parameters**

### **db** Db name

```
type: string
default: {}
optional: False
```

eval Evalue cutoff for blast to use

```
type: string
default: 1e-5
optional: True
```

group\_prefix OrthoMCL prefix for group names

```
type: string
default: g_
optional: True
```

#### host Db Host

```
type: localhost
default: {}
optional: True
```

input\_dir Input directory with compliant (read the manual) fasta files

```
type: string
default: {}
optional: False
```

# login Db username

```
type: string
default: None
optional: False
```

### mcl\_i mcl -i value

type: float default: 1.5 optional: True

### num\_threads Number of threads to use

type: integer default: 4 optional: True

# pass Db password

type: string default: None optional: False

# port Db port

type: integer default: 3306 optional: True

# prefix OrthoMCL prefix for the database tables

type: string default: ortho optional: True

#### vendor Db vendor

type: string default: mysql optional: True

### miscellaneous

```
Backend ruff
```

**Author** Mark Fiers

**Creation date** Tue Mar 29 16:34:19 2011

Modification date Wed Mar 30 06:02:01 2011

# 2.8.64 pregap

### **Pregap**

Run Pregap. Note that running phrap could be a part of this.

### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run no help defined

#### **Parameters**

cloning\_vector File containing the cloning vector

```
type: file
default: ''
optional: False
```

ecoli\_screenseq File containing ecoli screen sequences

```
type: file
default: ''
optional: False
```

input\_dir Directory with the input data

```
type: string
default: ''
optional: False
```

input\_pattern file name pattern

```
type: string
      default: "
      optional: False
quality_value_clip quality cutoff
      type: integer
      default: 10
      optional: True
repeat_masker_lib File with a repeatmasker library
      type: file
      default: "
      optional: False
sequencing_vector File containing the sequencing vector
      type: file
      default: ''
      optional: False
template the template pregap config file to use. if not defined, Moa tries ./files/pregap.config.
      type: file
      default: ./files/pregap.config.
      optional: True
vector_primerfile File with the vector primers
      type: file
      default: "
      optional: False
```

### miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.65 project

### Create a project

Placeholder for a Moa Project

#### **Commands**

run This template does not do anything - it is a project placeholder.

#### **Parameters**

#### miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Tue, 10 Jan 2012 14:54:39 +1300

Modification date Wed Nov 10 07:56:48 2010

### 2.8.66 reduce

### Execute a "reduce" ad-hoc analysis

Execute one command, on a number of input files.

#### **Commands**

run no help defined

#### **Filesets**

```
input "reduce" input files
output "reduce" output files
```

type: single category: output optional: True pattern: ./\*

#### **Parameters**

**process** The command to execute

type: string default: True optional: False

### miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Tue Mar 29 16:34:19 2011

Modification date Wed Mar 30 06:02:01 2011

### 2.8.67 sam2bam

### **Convert SAM to BAM using samtools**

Converts a FASTQ file to MAQ BFQ format.

#### **Commands**

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.

run no help defined

#### **Filesets**

input input SAM files

### output

type: map source: input category: output optional: {} pattern: ./\*.bam

### **Parameters**

#### miscellaneous

Backend gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.68 samtools pileup

Print the alignment in the pileup format.

#### **Commands**

```
clean Remove all job data, not the Moa job itselfrun run samtools pileup command
```

#### **Filesets**

fasta reference fasta file

```
type: single
category: prerequisite
optional: True
pattern: */*.fasta
```

### input bam or sam files

#### output

```
type: map
source: input
category: output
optional: {}
pattern: ./*.pileup
```

# output\_bam

```
type: map
source: input
category: output
optional: {}
pattern: ./*.sorted
```

### **Parameters**

cap\_mapQ\_at cap mapping quality at INT

type: integer default: 60

```
optional: True
extra_params any extra parameters
      type: string
      default: "
      optional: True
filter_read_bits filtering reads with bits in INT
      type: integer
      default: 1796
      optional: True
input_is_SAM the input is in SAM
      type: boolean
      default: False
      optional: True
num_haplotypes number of haplotypes in the sample (for -c/-g)
      type: integer
      default: 2
      optional: True
out_2nd_best output the 2nd best call and quality
      type: boolean
      default: False
      optional: True
out_GLFv3_format output in the GLFv3 format (suppressing -c/-i/-s)
```

type: boolean default: False optional: True

### out\_maq\_consensus output the maq consensus sequence

```
type: boolean
default: False
optional: True
```

# phred\_prob\_indel phred prob. of an indel in sequencing/prep. (for -c/-g)

```
type: integer
default: 40
optional: True
```

### print\_variants\_only print variants only (for -c)

```
type: boolean
default: False
optional: True
```

## **prior\_diff\_haplotypes** phred prob. of an indel in sequencing/prep. (for -c/-g)

```
type: float
default: 0.001
optional: True
```

# prior\_indel\_haplotypes number of haplotypes in the sample (for -c/-g)

```
type: float
default: 0.00015
optional: True
```

## **show\_lines\_indels** only show lines/consensus with indels

type: boolean default: False optional: True

# simple\_pileup\_format simple (yet incomplete) pileup format

type: boolean default: False optional: True

theta\_maq\_model number of haplotypes in the sample (for -c/-g)

type: float default: 0.85 optional: True

use\_SOAPsnp\_model use the SOAPsnp model for SNP calling

type: boolean default: False optional: True

### miscellaneous

Backend ruff

Author Yogini Idnani, Mark Fiers

**Creation date** Wed Dec 15 17:06:48 2010

Modification date unknown

### 2.8.69 sffinfo

#### sffinfo

Roche sffinfor tool - extract information from sff files

## Commands

clean Remove all job data, not the Moa job itself, note that this must be implemented by the template.

**run** Use the Roche sffinfo tool to extract reads, quality scores, flowgrams and accession ids from one or more sff files

#### **Filesets**

#### accession

type: map source: input category: output

```
optional: {}
      pattern: ./*.acc
flowgram
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./*.flow
input Sff input files
quality
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./*.qual
sequence
      type: map
      source: input
      category: output
      optional: {}
      pattern: ./*.reads
Parameters
accessions Output the accessions
      type: set
      default: T
      optional: True
flowgrams output the flowgrams
      type: set
      default: F
      optional: True
quality Output quality scores
      type: set
```

```
default: T
optional: True
```

### sequences Output the sequences

```
type: set
default: T
optional: True
```

# untrimmed output untrimmed sequences & qualities

```
type: set
default: F
optional: True
```

#### miscellaneous

**Backend** gnumake

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

# 2.8.70 simple

# Execute a "simple" ad hoc analysis

Execute one command, No in or output files are tracked by Moa.

### **Commands**

run no help defined

#### **Parameters**

**process** The command to execute

type: string default: True optional: False

### miscellaneous

Backend ruff

**Author** Mark Fiers

Creation date Tue Mar 29 16:34:19 2011

Modification date Wed Mar 30 06:02:01 2011

# 2.8.71 smalt\_pe

Run SMALT on an set of input files (query) vs a database index.

#### **Commands**

clean Remove all job data, not the Moa job itselfrun Execute SMALT with with paired-end fastq

#### **Filesets**

**db** The (basename of the) smalt database to use.

type: single

category: prerequisite

optional: False

pattern: ../10.smaltdb/db

fasta reference fasta file

type: single

category: prerequisite

optional: False pattern: \*.fasta

fq\_forward\_input fastq input files directory - forward

fq\_reverse\_input fastq input files directory - reverse

type: map

source: fq\_forward\_input

category: input optional: True pattern: \*/\*\_2.fq

**output** output BAM file (automatically converted & filtered for reads that to not map)

```
type: map
source: fq_forward_input
category: output
optional: {}
pattern: ./*.sam
```

### **Parameters**

extra\_params extra parameters to feed to smalt

```
type: string
default: ''
optional: True
```

max\_insertsize Maximum allowed insertsize

```
type: integer
default: 250
optional: True
```

min insertsize Minimum allowed insertsize

```
type: integer
default: 1
optional: True
```

output\_format output format (sam or samsoft)

```
type: {}
default: sam
optional: True
```

pairtype pair type (pe: fr/illumina short; mp: rf/illumina mate pairs or pp: ff

```
type: {}
default: pe
```

```
optional: True
```

#### threads No threads to use

```
type: int
default: 4
optional: True
```

#### miscellaneous

**Backend** ruff

**Author** Mark Fiers

Creation date Tue, 27 Mar 2012 10:05:40 +1300

**Modification date** Tue, 27 Mar 2012 10:31:09 +1300

#### 2.8.72 smaltdb

#### Smalt index builder

Builds a smalt index from a reference sequence

### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** Create the smalt index

### **Filesets**

input Input fasta file for the smalt database

```
type: single
category: input
optional: False
pattern: */*.fasta
```

# output database name to create

```
type: single
category: output
optional: {}
pattern: db
```

### **Parameters**

word\_length word length

type: int default: 10 optional: True

word\_spacing word spacing

type: int default: 6 optional: True

### miscellaneous

Backend ruff

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

**Modification date** Wed Dec 09 07:56:48 2010

# 2.8.73 soapdenovo\_pe

Run Soapdenovo

#### **Commands**

clean Remove all job data

run Execute soapdenovo in paired-end mode

#### **Filesets**

fq\_forward fastq input files directory - forward

 $fq\_reverse$  fastq input files directory - reverse

type: map

source: fq\_forward category: input optional: True pattern: \*/\*\_2.fq

### output soap denovo output file

```
type: single
category: output
optional: True
pattern: {}
```

#### **Parameters**

avg\_insert library insert size

```
type: integer
default: 200
optional: {}
```

**executable** which executable to use (SOAPdenovo-127mer, SOAPdenovo-31mer or SOAPdenovo-63mer)

```
type: {}
default: SOAPdenovo-31mer
optional: True
```

kmer kmer size

type: integer default: 31 optional: True

**skip\_config\_file** skip automatic config file generation - if you skip this, make sure that you have a soap.config configuration file in the current directory

type: boolean default: False optional: True

threads no threads to use

type: integer

```
default: 8
optional: True
```

### miscellaneous

```
Backend ruff
Author Mark Fiers
Creation date Mon, 21 Nov 2011 12:47:16
```

Modification date Mon, 21 Nov 2011 12:47:22

### 2.8.74 statsidx

Retrieve and print stats from BAM file to an index file

### **Commands**

```
clean Remove all job data, not the Moa job itselfrun run samtools idxstats
```

#### **Filesets**

```
input bam input files directory - forward files
output
```

```
type: map
source: input
category: output
optional: {}
pattern: ./*.index
```

#### **Parameters**

#### miscellaneous

```
Backend ruff
Author Yogini Idnani, Mark Fiers
Creation date Wed Dec 08 17:06:48 2010
Modification date unknown
```

## 2.8.75 sync

## Sync directories

Create this directory in sync with another directory

## **Commands**

```
run Sync!
```

#### **Parameters**

ignore ignore these names (space separated list)

```
type: {}
default: "
optional: True
```

**original** The local directory to use as a source. If the target (based on what is in the source) does not exists, this directory is copied. If the target exists - only the configuration is copied, and all directory contents are left alone. If this parameter is omitted, the directory with the most recently changed moa configuration.

```
type: string
default: {}
optional: True
```

recursive copy the jobs/config recursively

```
type: boolean
default: False
optional: True
```

source The directory to keep in sync with

```
type: string
default: {}
optional: False
```

2.8. Templates

#### miscellaneous

```
Backend ruff
Author Mark Fiers
Creation date Thu, 30 Jun 2011 21:26:19
Modification date Thu, 30 Jun 2011 21:25:53
```

#### 2.8.76 unittest

Template used in testing - has no other purpose

#### **Commands**

```
clean Remove all job data
prepare prepare for the unittest
run Prepare & Run
run delegates execution to: prepare, run2
run2 actually run
```

#### **Filesets**

```
input_1 Input file set 1
input_2 Input file set 2

type: map
    source: input_1
    category: input
    optional: {}
    pattern: in2/*_2.txt
```

## output output files

```
type: map
source: input_1
category: output
optional: {}
pattern: ./*.out
```

#### **Parameters**

test\_string Test string values

```
type: string
default: {}
optional: True
```

#### miscellaneous

```
Backend ruff
```

Author Yogini Idnani, Mark Fiers

**Creation date** Wed Nov 25 17:06:48 2010

Modification date unknown

## 2.8.77 varscan

#### Varscan

Run VARSCAN to detect snps

#### **Commands**

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** *no help defined* 

#### **Parameters**

extra\_params location of varscan.pl, defaults to /usr/lib/perl5/site\_perl/5.8.8/varscan.pl

```
type: string
default: "
optional: True
```

input\_file Varscan input alignments file

```
type: file
default: "
optional: True
```

output\_name Base name of the output files

2.8. Templates

```
type: string
default: out
optional: True
```

perl\_file the varscan (perl) executable

```
type: file
default: "
optional: True
```

#### miscellaneous

```
Backend gnumake
```

**Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

Modification date Wed Nov 10 07:56:48 2010

## 2.8.78 vpcr

#### **VPCR**

Virtual PCR, based on Bowtie

## Commands

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** Predict the fragments that would be generated by a PCR

#### **Parameters**

bowtie\_db Location of the bowtie database used for the vpcr

```
type: file
default: "
optional: True
```

insert\_max maximum insert size for a vpcr fragment

type: integer default: 10000 optional: True

## insert\_min minimal insert size for a fragment

```
type: integer
default: 10
optional: True
```

## primer\_1 First primer to use

```
type: string
default: "
optional: False
```

## primer\_2 Second primer to use

```
type: string
default: ''
optional: False
```

#### miscellaneous

```
Backend gnumake
Author Mark Fiers
Creation date Wed Nov 10 07:56:48 2010
```

Modification date Wed Nov 10 07:56:48 2010

## 2.8.79 vpcr\_list

Virtual PCR, based on Bowtie

#### Commands

**clean** Remove all job data, not the Moa job itself, note that this must be implemented by the template. **run** Predict the fragments that would be generated by a PCR

#### **Parameters**

bowtie\_db Location of the bowtie database used for the vpcr

type: file

2.8. Templates

```
default: ''
optional: False
```

insert\_max maximum insert size for a vpcr fragment

```
type: integer
default: 10000
optional: True
```

insert\_min minimal insert size for a fragment

```
type: integer
default: 10
optional: True
```

primer\_list List of primers to check

```
type: file
default: {}
optional: False
```

#### miscellaneous

**Backend** gnumake **Author** Mark Fiers

**Creation date** Wed Nov 10 07:56:48 2010

**Modification date** Wed Nov 10 07:56:48 2010

## 2.8.80 wget

## wget

Use WGET to download files. This template has two modi, one is set wget\_mode to mirror data, in which case both wget\_url and wget\_pattern (default \*) are used. The other modus is wget\_mode=get, when one file defined by wget\_url is downloaded. In the mirror mode it is possible to download only those files that are newer as the files already downloaded by using the wget\_timestamp parameter

#### **Commands**

run Download

#### **Parameters**

pass Password for the remote site (note - this is not very safe, the password will be stored in plan text

```
type: password
default: "
optional: True
```

url The url of the file to download

```
type: string
default: {}
optional: False
```

user Username for the remote site

```
type: string
default: "
optional: True
```

#### miscellaneous

```
Backend ruff
```

**Author** Mark Fiers

Creation date Thu, 02 Jun 2011 10:22:31 +1200

Modification date Thu, 02 Jun 2011 10:22:53 +1200

## 2.9 Moa API

## 2.9.1 moa.actor

'Simple' wrapper around subprocess to execute code

```
moa.actor.getLastStderr (job)
Get the last stderr

moa.actor.getLastStdout (job)
Get the last stdout

moa.actor.getRecentOutDir (job)
Return the most recent output directory
```

```
moa.actor.simpleRunner (wd, cl, conf={}, **kwargs)

Don't think - just run - here & now
```

what does this function do? - put env in the environment - Execute the commandline (in cl) - store stdout & stderr in log files - return the rc

#### 2.9.2 moa.commands

Handle Moa commands (i.e. anything that you can run as moa COMMAND on the commandline

## 2.9.3 moa.job

```
{f class} moa. job. {f Job} (wd)
```

Class defining a single job

Note - in the moa system, there can be only one current job - many operations try to access the job in sysConf

```
>>> wd = tempfile.mkdtemp()
>>> job = Job(wd)
>>> assert(isinstance(job, Job))
>>> assert(job.template.name == 'nojob')
```

#### checkCommands (command)

Check command, and rearrange if there are delegates.

```
>>> job = newTestJob('unittest')
>>> assert(job.template.commands.run.delegate == ['prepare', 'run2'])
>>> assert(job.checkCommands('run2') == ['run2'])
>>> assert(job.checkCommands('run') == ['prepare', 'run2'])
>>> assert(job.checkCommands('prepare') == ['prepare'])
```

**Parameters commands** (*list of strings*) – The list of commands to check

**Returns** The checked list of commands

**Return type** list of strings

#### checkConfDir()

Check if the configuration directory exists. If not create it.

```
>>> job = newTestJob('unittest')
>>> confdir = os.path.join(job.wd, '.moa')
>>> assert(os.path.exists(confdir))
>>> import shutil
>>> shutil.rmtree(confdir)
>>> assert(os.path.exists(confdir) == False)
>>> job.checkConfDir()
>>> assert(os.path.exists(confdir))
```

## ${\tt defineCommands}\ (commandparser)$

Register template commands with the argparser

```
defineOptions (parser)
```

Set command line options - deferred to the backend - PER COMMAND

```
>>> job = newTestJob('unittest')
     >>> import optparse
     >>> parser = optparse.OptionParser()
     >>> job.defineOptions(parser)
execute (job, args, **kwargs)
     Execute command in the context of this job. Execution is always deferred to the backend
     #Note: Uncertain how to test verbose & silent
         Parameters
             • verbose (Boolean) – output lots of data
             • silent (Boolean) – output nothing
finishExecute()
     Finish the run!
getFiles()
     Return all moa files - i.e. all files crucial to this job.
hasCommand (command)
     Check if this job defines a certain command
      Warning: THIS METHOD DOES NOT WORK PROPERLY YET
     >>> job = newTestJob('unittest')
     >>> assert (job.hasCommand('run'))
     >>> assert (not job.hasCommand('dummy'))
initialize()
     Initialize a new job in the current wd
isMoa()
     Check if this is a Moa directory - Currently, this needs to be overridden #weird; uncertain if
     this ever gets called
loadBackend()
     load the backend
loadTemplate()
     Load the template for this job, based on what configuration can be found
loadTemplateMeta()
     Load the template meta data for this job, based on what configuration can be found
prepareExecute()
     Give this job a chance to prepare for execution.
```

>>> os.unlink(templateFile)
>>> assert(not os.path.exists(templateFile))

>>> templateFile = os.path.join(job.confDir, 'template.d', 'unittest.jinja2')

2.9. Moa API 149

Reload the template into the local .moa/template.d directory

>>> assert (os.path.exists(templateFile))

>>> job = newTestJob('unittest')

refreshTemplate()

```
>>> job.refreshTemplate()
>>> assert (os.path.exists(templateFile))

run_hook (hook, **kwargs)
    Shortcut to run a job plugin hook

setTemplate (name, provider=None)
    Set a new template for this job

>>> job = newTestJob('unittest')
>>> job.setTemplate('adhoc')
>>> afile = os.path.join(job.confDir, 'template.d', 'adhoc.mk')
>>> assert (os.path.exists(afile))

moa.job.newJob (wd, template, title, parameters=[], provider=None)
Create a new job in the wd and return the proper job object currently only makefile jobs are supported - later we'll scan the template, and instantiate the proper job type
```

```
>>> wd = tempfile.mkdtemp()
>>> job = newJob(wd, template='blast', title='test')
>>> assert(isinstance(job, Job))
>>> assert(job.template.name == 'blast')
>>> assert(job.conf.title == 'test')
```

#### **Parameters**

- wd Directory to create this job in, note that this directory must already exists
- **template** (*String*) Template name for this job
- parameters (list of (key, value) tuples) A list of parameters to set for this job

**Return type** instance of moa.job.Job

```
moa.job.newTestJob (template, title='Test job', provider=None)
```

for testing purposes - creates a temporary directory and uses that to instantiate a job. This function returns the job object created

```
>>> job = newTestJob(template = 'adhoc', title='test title')
>>> assert(isinstance(job, Job))
>>> assert(os.path.exists(job.wd))
>>> assert(job.conf.title == 'test title')
>>> assert(os.path.exists(os.path.join(job.wd, '.moa')))
>>> assert(os.path.exists(os.path.join(job.wd, '.moa', 'template')))
>>> assert(job.template.name == 'adhoc')
```

Returns the created job

Return type instance of moa.job.Job

## 2.9.4 moa.jobConf

moa job configuration

## ${f class} \; {f moa.jobConf.JobConf} \; (job)$

to distinguish between attributes of this object & proper job configuration parameters

#### doNotCheck = None

these fields are not be type-checked

#### doNotSave = None

these fields are not to be saved

#### getRendered(key)

Get the rendered value of this key

#### isEmpty()

Check if the config is empty is empty

#### isPrivate(k)

Is this a private variable? can be locally defined or in the template definition

#### keys()

return a dict with all known parameters and values, either defined in the job configuration of the template

#### load (confFile, delta=None)

Load a configuration file

**Parameters delta** – if a value appears to be a relative path, try to correct for this. Currently this only works for files that exist. i.e.

## private = None

these fields are private (i.e. not to be displayed by default)

#### save()

Save the conf to disk

#### setRecursiveVar(k, v)

Register a recursive variable

#### 2.9.5 moa.sysConf

Store Moa wide configuration

#### 2.9.6 moa.ui

communicate information to the user

#### 2.9.7 moa.utils

A set of random utilities used by Moa

#### moa.utils.deprecated(func)

Decorator function to flag a function as deprecated

**Parameters func** – any function

```
moa.utils.flog(f)
```

A simple logger - uses the moa.logger code to log the calling function. Use as a decorator:

```
@moa.utils.flog
def any_function(*args);
```

This is for debugging purposes (obviously)

Parameters func – Any python function

```
moa.utils.getMoaBase()
```

Return MOABASE - the directory where Moa is installed. This function also sets an environment variable *MOABASE* 

```
>>> d = getMoaBase()
>>> assert(os.path.isdir(d))
>>> assert(os.path.isfile(os.path.join(d, 'README')))
>>> assert(os.path.isdir(os.path.join(d, 'lib')))
```

**Return type** string (path)

```
moa.utils.getProcessInfo(pid)
```

Return some info on a process

```
moa.utils.moaDirOrExit(job)
```

Check if the job contains a proper Moa job, if not, exit with an error message and a non-zero exit code.

Parameters job - An instance of moa.job.Job

```
moa.utils.simple_decorator(decorator)
```

This decorator can be used to turn simple functions into well-behaved decorators, so long as the decorators are fairly simple. If a decorator expects a function and returns a function (no descriptors), and if it doesn't modify function attributes or docstring, then it is eligible to use this. Simply apply @simple\_decorator to your decorator and it will automatically preserve the docstring and function attributes of functions to which it is applied.

Note; I got this code from somehwere, but forgot where exactly. This seems the most likely source:

http://svn.navi.cx/misc/trunk/djblets/djblets/util/decorators.py

#### 2.9.8 moa.template

## moa.template

Store information on a template. This module is also responsible for retrieving template information.

```
moa.template.initTemplate(*args, **kwargs)
```

```
moa.template.installTemplate(wd, tName, provider=None)
```

Initialize the template - this means - try to figure out where the template came from & copy the template files into *job/.moa/template & job/.moa/template.d/extra*.

Currently all templates come from the moa repository. In the future, multiple sources must be possible

```
>>> import tempfile
>>> wd = tempfile.mkdtemp()
>>> installTemplate(wd, 'adhoc')
>>> templateFile = os.path.join(wd, '.moa', 'template')
>>> adhocFile = os.path.join(wd, '.moa', 'template.d', 'adhoc.mk')
>>> assert(os.path.exists(templateFile))
>>> assert(os.path.exists(adhocFile))
moa.template.refresh(wd)
```

Refresh the template - try to find out what the template is from {{wd}}/.moa/template.d/meta. If that doesn't work, revert to the default template. If default is not specified - exit with an error

```
>>> import tempfile
>>> wd = tempfile.mkdtemp()
>>> installTemplate(wd, 'adhoc')
>>> templateFile = os.path.join(wd, '.moa', 'template')
>>> adhocFile = os.path.join(wd, '.moa', 'template.d', 'adhoc.mk')
>>> os.unlink(adhocFile)
>>> os.unlink(templateFile)
>>> assert(not os.path.exists(templateFile))
>>> refresh(wd)
>>> assert(os.path.exists(templateFile))
>>> assert(os.path.exists(adhocFile))
```

#### moa.template.template

Store information on a template. This module is also responsible for retrieving template information.

```
class moa.template.template.Template(templateFile)
    Template extends Yaco
    getRaw()
```

Return a Yaco representation of the yaml-template, without any of this Template processing. This is really useful when processing a template that needs to be written back to disk

```
>>> import moa.job
>>> job = moa.job.newTestJob(template='adhoc')
>>> raw = job.template.getRaw()
>>> assert(isinstance(raw, Yaco.Yaco))
>>> assert(raw.has_key('parameters'))
```

## 2.9.9 moa.template.provider

#### moa.provider.core

Provides templates from the Moa package.

#### 2.9.10 moa.backend

## Ruff

Ruffus (and Jinja) Backend

#### members

## 2.9.11 moa.plugin

#### adhoc - create jobs from adhoc bash code

```
moa.plugin.adhoc.createAdhoc(job)
Creates an adhoc job.
moa.plugin.adhoc.createMap(job, args)
create an adhoc moa 'map' job
```

Moa will query the user for process, input & output files. An example session

```
moa.plugin.adhoc.createReduce (job) Create a 'reduce' adhoc job.
```

There are a number of ways this command can be used:

```
$ moa reduce -t 'a title' -- echo 'define a command'
```

Anything after – will be the executable command. If omitted, Moa will query the user for a command.

Moa will also query the user for input & output files. An example session:

```
$ moa map -t 'something intelligent'
process:
> echo 'processing {{ input }} {{ output }}'
input:
> ../10.input/*.txt
output:
> ./*.out
```

Assuming you have a number of text files in the ../10/input/ directory, you will see, upon running:

```
processing ../10.input/test.01.txt ./test.01.out processing ../10.input/test.02.txt ./test.02.out processing ../10.input/test.03.txt ./test.03.out ...
```

```
moa.plugin.adhoc.exclamate(job, args)
```

Create a 'simple' job from the last command issued.

Set the *process* parameter to the last issued command. If a moa job exists in the current directory, then the *process* parameter is set without questions. (even if the Moa job in question does not use the *process* parameter). If no moa job exists, a *simple* job is created first.

Note: This works only when using bash and if moainit is sourced properly. moainit defines a bash function \_moa\_prompt that is called every time a command is issued (using \$PROMPT\_COMMAND). The \_moa\_prompt function takes the last command from the bash history and stores it in ~/.config/moa/last.command. Additionally, the \_moa\_prompt function stores all commands issued in a Moa directory in .moa/local\_bash\_history.

```
moa.plugin.adhoc.exclamateInJob (job, args, command)
```

Reuse the last issued command: set it as the 'process' parameters in the current job

```
moa.plugin.adhoc.exclamateNoJob (job, args, command)

Create a "simple" job & set the last command to the 'process' parameter

moa.plugin.adhoc.simple (job, args)

Create a 'simple' adhoc job.
```

Simple meaning that no in or output files are tracked. Moa will query you for a command to execute (the *process* parameter).

## 2.9.12 Yaco

Yaco provides a *dict* like structure that can be serialized to & from yaml. Yaco objects behave as dictionaries but also allow attribute access (loosely based on this 'recipe < http://code.activestate.com/recipes/473786/>'\_). Sublevel dictionaries are automatically converted to Yaco objects, allowing sublevel attribute access, for example:

```
>>> x = Yaco()
>>> x.test = 1
>>> x.sub.test = 2
>>> x.sub.test
```

Note that sub-dictionaries do not need to be initialized. This has as a consequence that requesting uninitialized items automatically return an empty Yaco object (inherited from a dictionary).

Yaco can be found in the Python package index and is also part of the Moa source distribution

## **Autogenerating keys**

An important feature (or annoyance) of Yaco is the auto generation of keys that are not present (yet). For example:

```
>>> x = Yaco()
>>> x.a.b.c.d = 1
>>> assert(x.a.b.c.d == 1)
```

works - a, b and c are assumed to be Yaco dictionaries and d is give value 1. This makes populating data structures easy.

It might also generate some confusion when querying for keys in the Yaco structure - if a key does not exists, it automatically comes back as an empy *dict* or *Yaco* object (renders as {}). This means that if it is easy to check if a certain 'branch' of a Yaco datastructure exists:

```
>>> x = Yaco()
>>> assert (not x.a.b)
```

but now the following works as well:

```
>>> assert(x.has_key('a'))
>>> assert(x.a.has_key('b'))
```

So, a safe way to test a data structure, without introducing extra branches is:

```
>>> x = Yaco()
>>> assert(not x.has_key('a'))
```

Todo: Need to find a more elegant way of testing without introducing data structures

```
class Yaco.Yaco (data={})
     Rather loosely based on http://code.activestate.com/recipes/473786/ (r1)
     >>> v= Yaco()
     >>> v.a = 1
     >>> assert (v.a == 1)
     >>> assert(v['a'] == 1)
     >>> v= Yaco({'a':1})
     >>> assert(v.a == 1)
     >>> assert(v['a'] == 1)
     get_data()
         Prepare & parse data for export
         >>> y = Yaco()
         >>> y.a = 1
         >>> y.b = 2
         >>> y._c = 3
         >>> assert (y._c == 3)
         >>> d = y.get_data()
         >>> assert (d.has_key('a') == True)
         >>> assert (d.has_key('b') == True)
         >>> assert (d.has_key('_c') == False)
         >>> y._private = ['b']
         >>> d = y.get_data()
         >>> assert (d.has_key('a') == True)
         >>> assert (d.has_key('b') == False)
         >>> assert (d.has key(' c') == False)
     load (from_file)
         Load this dict from_file
         >>> import yaml
         >>> import tempfile
         >>> tf = tempfile.NamedTemporaryFile(delete=False)
         >>> tf.write(yaml.dump({'a' : [1,2,3, [1,2,3, {'d' : 4}]], 'b': 4, 'c': '5'}))
         >>> tf.close()
         >>> y = Yaco()
         >>> y.load(tf.name)
         >>> assert (y.a[3][3].d == 4)
    pretty()
         Return data as a pprint.pformatted string
     save (to_file, doNotSave= | )
     simple()
         return a simplified representation of this Yaco struct - remove Yaco from the equation - and
         all object reference. Leave only bool, float, str, lists, tuples and dicts
         >>> x = Yaco()
         >>> x.y.z = 1
         >>> assert (isinstance (x.y, Yaco))
         >>> s = x.simple()
         >>> assert(s['y']['z'] == 1)
         >>> assert(isinstance(s['y'], dict))
         >>> assert (not isinstance (s['y'], Yaco))
```

#### update (data)

```
>>> v = Yaco({'a' : [1,2,3,{'b' : 12}]})
>>> assert(v.a[3].b == 12)

>>> v = Yaco({'a' : [1,2,3,[1,{'b' : 12}]]})
>>> assert(v.a[3][1].b == 12)
```

#### 2.9.13 fist

#### **Filesets**

Handle & manipulate sets of files

This module aims at providing classes to handle and manipulate sets of files. Two simple examples are a simple set containing one file (fist.fistSingle) or a *glob* based set of files (fist.fistFileset). A more complicated example is fistMapset that maps another fileset based on a pattern.

Each fileset inherits from *list* - hence fist filesets behave as lists.

Future work should allow the definition of remote filesets (for example over http or ssh).

Each fist class is istantiated with a url defining the file(set). In the case of fist.fistFileset this url contains a globbing characters:

```
fs = fist.fistFileset('/tmp/*,txt')
```

This fileset object contains a list with all \*.txt files in /tmp. Subsequently it is possible to map this set

```
class fist.fistCore(url, context=None)
```

Core class for all fist classes

```
resolve()
```

This function needs to be overridden context

```
class fist.fistFileset (url, context=None)
```

Most basic set of files - handle a set of files described by a single URI with wildcards, for example:

```
* '*.txt'
* '../*.txt'
* 'file:///home/name/data/*.txt'
>>> f = fistFileset('*.txt')
>>> assert (f.path=='.')
>>> assert(f.glob=='*.txt')
>>> assert (f.path=='.')
>>> assert(f.glob=='*.txt')
>>> f = fistFileset('/tmp')
>>> assert(f.path=='/tmp')
>>> assert(f.glob==' *')
>>> f = fistFileset('/tmp/*.txt')
>>> assert (f.path=='/tmp')
>>> assert(f.glob=='*.txt')
>>> f = fistFileset('../*.txt')
>>> assert(f.path=='...')
>>> assert(f.glob=='*.txt')
```

```
>>> f = fistFileset(os.path.join(wd, 'in', '*.txt'))
    >>> f.resolve()
    >>> assert (len(f) == 100)
    >>> f = fistFileset(os.path.join(wd, 'in', 'in1*.txt'))
    >>> f.resolve()
    >>> assert (len(f) == 10)
    >>> f = fistFileset('~/*')
    >>> f.resolve()
    >>> assert (len(f) > 0)
class fist.fistMapset (url, context=None)
    fistMapset
    Map set - map a fileset based on a target uri
    >>> f = fistFileset(os.path.join(wd, 'in', '*'))
    >>> f.resolve()
    >>> assert(len(f) == 100)
    >>> ##
    >>> ## Null mapping
    >>> m = fistMapset('*/*')
    >>> m.resolve(f)
    >>> assert (len (m) == 100)
    >>> assert (os.path.join(wd, 'in/in18.txt') in m)
    >>> ##
    >>> ## simple folder mapping
    >>> ##
    >>> m = fistMapset('out/*')
    >>> m.resolve(f)
    >>> assert (len (m) == 100)
    >>> assert('out/in18.txt' in m)
    >>> ##
    >>> ## simple folder mapping
    >>> ##
    >>> m = fistMapset('./*')
    >>> m.resolve(f)
    >>> assert (len (m) == 100)
    >>> assert('./in18.txt' in m)
    >>> ##
    >>> ## simple folder & mapping & extension append
    >>> ##
    >>> m = fistMapset('out/*.out')
    >>> m.resolve(f)
    >>> assert (len (m) == 100)
    >>> assert('out/in18.txt.out' in m)
    >>> ##
    >>> ## New from fileset - now with a pattern defining the extension
    >>> f = fistFileset(os.path.join(wd, 'in', '*.txt'))
    >>> f.resolve()
    >>> ## extension mapping
    >>> ##
    >>> m = fistMapset('out/*.out')
    >>> m.resolve(f)
    >>> assert (len (m) == 100)
    >>> assert('out/in18.out' in m)
```

```
>>> ##
     >>> ## New from fileset - now with a pattern defining file glob &
     >>> ##
     >>> f = fistFileset(os.path.join(wd, 'in', 'in*.txt'))
     >>> f.resolve()
     >>> ## more complex filename mapping
     >>> ##
     >>> m = fistMapset('out/test*.out')
     >>> m.resolve(f)
     >>> assert (len (m) == 100)
     >>> assert('out/test18.out' in m)
     >>> ## mapping keeping the extension the same
    >>> m = fistMapset('out/test*.txt')
     >>> m.resolve(f)
     >>> assert (len (m) == 100)
    >>> assert('out/test18.txt' in m)
     resolve (mapFrom)
         Resolve the mapped set based on a input fileSet
     resolver (mapFrom, list)
         map all files in the incoming list
class fist.fistSingle(url, context=None)
     Represents a single file
     init()
         Assuming the url is a single file
```

CHAPTER

**THREE** 

# **MORE INFORMATION**

**CHAPTER** 

**FOUR** 

# **INDICES AND TABLES**

- genindex
- modindex
- search

## **PYTHON MODULE INDEX**

```
f
fist, 157
m
moa.actor, 147
moa.backend.ruff, 153
moa.commands, 148
moa.job, 148
moa.jobConf, 150
moa.plugin.adhoc, 154
moa.sysConf, 151
moa.template, 152
moa.template.provider.core, 153
moa.template.template, 153
moa.ui, 151
moa.utils, 151
У
Yaco, 155
```

## **INDEX**

C checkCommands() (moa.job.Job method), 148 checkConfDir() (moa.job.Job method), 148 createAdhoc() (in module moa.plugin.adhoc), 154 createMap() (in module moa.plugin.adhoc), 154 createReduce() (in module moa.plugin.adhoc), 154	getRecentOutDir() (in module moa.actor), 147 getRendered() (moa.jobConf.JobConf method), 151  H hasCommand() (moa.job.Job method), 149
D	I
defineCommands() (moa.job.Job method), 148 defineOptions() (moa.job.Job method), 148 deprecated() (in module moa.utils), 151 doNotCheck (moa.jobConf.JobConf attribute), 151 doNotSave (moa.jobConf.JobConf attribute), 151	init() (fist.fistSingle method), 159 initialize() (moa.job.Job method), 149 initTemplate() (in module moa.template), 152 installTemplate() (in module moa.template), 152 isEmpty() (moa.jobConf.JobConf method), 151 isMoa() (moa.job.Job method), 149 isPrivate() (moa.jobConf.JobConf method), 151
E	J
exclamate() (in module moa.plugin.adhoc), 154 exclamateInJob() (in module moa.plugin.adhoc), 154	Job (class in moa.job), 148 JobConf (class in moa.jobConf), 150
exclamateNoJob() (in module moa.plugin.adhoc), 154	K
1 14	
execute() (moa.job.Job method), 149	keys() (moa.jobConf.JobConf method), 151
	keys() (moa.jobConf.JobConf method), 151
execute() (moa.job.Job method), 149  F finishExecute() (moa.job.Job method), 149 fist (module), 157 fistCore (class in fist), 157 fistFileset (class in fist), 157 fistMapset (class in fist), 158	L load() (moa.jobConf.JobConf method), 151 load() (Yaco.Yaco method), 156 loadBackend() (moa.job.Job method), 149 loadTemplate() (moa.job.Job method), 149 loadTemplateMeta() (moa.job.Job method), 149
execute() (moa.job.Job method), 149  F  finishExecute() (moa.job.Job method), 149 fist (module), 157 fistCore (class in fist), 157 fistFileset (class in fist), 157	L load() (moa.jobConf.JobConf method), 151 load() (Yaco.Yaco method), 156 loadBackend() (moa.job.Job method), 149 loadTemplate() (moa.job.Job method), 149 loadTemplateMeta() (moa.job.Job method), 149  M
execute() (moa.job.Job method), 149  F  finishExecute() (moa.job.Job method), 149  fist (module), 157  fistCore (class in fist), 157  fistFileset (class in fist), 157  fistMapset (class in fist), 158  fistSingle (class in fist), 159	L load() (moa.jobConf.JobConf method), 151 load() (Yaco.Yaco method), 156 loadBackend() (moa.job.Job method), 149 loadTemplate() (moa.job.Job method), 149 loadTemplateMeta() (moa.job.Job method), 149

```
moa.ui (module), 151
moa.utils (module), 151
moaDirOrExit() (in module moa.utils), 152
Ν
newJob() (in module moa.job), 150
newTestJob() (in module moa.job), 150
Р
prepareExecute() (moa.job.Job method), 149
pretty() (Yaco. Yaco method), 156
private (moa.jobConf.JobConf attribute), 151
R
refresh() (in module moa.template), 153
refreshTemplate() (moa.job.Job method), 149
resolve() (fist.fistCore method), 157
resolve() (fist.fistMapset method), 159
resolver() (fist.fistMapset method), 159
run_hook() (moa.job.Job method), 150
S
save() (moa.jobConf.JobConf method), 151
save() (Yaco. Yaco method), 156
setRecursiveVar()
                          (moa.jobConf.JobConf
        method), 151
setTemplate() (moa.job.Job method), 150
simple() (in module moa.plugin.adhoc), 155
simple() (Yaco. Yaco method), 156
simple_decorator() (in module moa.utils), 152
simpleRunner() (in module moa.actor), 147
Τ
Template (class in moa.template.template), 153
U
update() (Yaco. Yaco method), 156
Υ
Yaco (class in Yaco), 155
Yaco (module), 155
```

168 Index