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# **BioMAJ Documentation**

*Release 3.0*

**Olivier Sallou**

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Getting Started Documentation:



For a very basic setup, you can configure a `docker-compose.yml` file to use with `docker`, which is especially helpful when you are testing out BioMAJ.

### 1.1 Docker

```
1 version: '2'
2 services:
3   biomaj:
4     image: osallou/biomaj-docker
5     links:
6       - mongodb:biomaj-mongodb
7     volumes:
8       - ./data:/var/lib/biomaj
9
10  mongodb:
11    image: mongo
```

This configuration file defines a simple MongoDB instance which is used for backend storage by BioMAJ, as well as the BioMAJ instance itself. Line 8 denotes that a folder named `data` in the current directory will be mounted into the volume as storage. Any files downloaded by BioMAJ will appear in this directory.

Running the `--help` command can be done easily:

```
$ docker-compose run --rm biomaj --help
```

### 1.2 Simple Configuration

Once you've reached this point, you're ready to start configuring BioMAJ to download datasets for you. Configuration files should go instead a folder `conf` inside the `data` folder in your current directory. As an example, we will use this simple ALU configuration file:

```
1 [GENERAL]
2 # Database name/description
3 db.fullname="alu.n : alu repeat element. alu.a : translation of alu.n repeats"
4 # The short name for the database
5 db.name=alu
6 # Database type. Some common values include genome, nucleic, nucleic_protein, protein,
7 ↪ other
8 db.type=nucleic_protein
9 # Base directory to download to download temp files to
10 offline.dir.name=offline/ncbi/blast/alu_tmp
11 # Base directory to download to
12 dir.version=ncbi/blast/alu
13 # Update frequency
14 frequency.update=0
15 # Number of threads used during downloading
16 files.num.threads=1
17
18 # Protocol, common values include ftp, http
19 protocol=ftp
20 # The FQDN of the server you with to connect to
21 server=ftp.ncbi.nih.gov
22 # And the directory on that server
23 remote.dir=/blast/db/FASTA/
24 # The files to find in that page of the remote server.
25 remote.files=^alu.*\.gz$
26
27 # BioMAJ can automatically extract the version number from a release
28 # document. This will be covered in another section.
29 release.file=
30 release.regexp=
31 release.file.compressed=
32
33 #Uncomment if you don't want to extract the data files.
34 #no.extract=true
35
36 # ?
37 local.files=^alu\.(a|n).*
38
39 ## Post Process ## The files should be located in the projectfiles/process directory
40 db.post.process=
41
42 ### Deployment ###
43 keep.old.version=1
```

The file can be broken down into a couple of sections:

- Metadata (lines 1-15)
- Remote Source (17-24)
- Release Information (26-30)
- Other

The metadata consists of things like where data should be stored, and how to name it. The remote source describes where data is to be fetched from, release information we will see in another example, and then there are a few extra, miscellaneous options shown in this example config.

If you have copied the `alu.properties` file into `./data/conf/alu.properties`, you are ready to download this database:



```
$ docker-compose run --rm biomaj --bank alu --update
2016-08-24 21:43:15,276 INFO [root][MainThread] Log file: /var/lib/biomaj/log/alu/
↪1472074995.28/alu.log
Log file: /var/lib/biomaj/log/alu/1472074995.28/alu.log
...
```

This command should complete successfully, and you will have some more files in `./data/`:

```
$ find data
data/conf/alu.properties
data/data/ncbi/blast/alu/alu-2003-11-26/flat/alu.a
data/data/ncbi/blast/alu/alu-2003-11-26/flat/alu.n
data/cache/files_1472074995.29
data/log/alu/1472074995.28/alu.log
```

The `data/data` directories contain your downloaded files. Additionally a cache file exists and a job run log is contains data about what occurred during the download and processing. Note that the files that appear are `alu.a` and `alu.n`, instead of `alu.a.gz` and `alu.n.gz`. By having the option `no.extract=true` commented out on line 33, BioMAJ automatically extracted the data for us.

The `--status` command will allow you to see the status of various databases you have downloaded.

```
$ docker-compose run --rm biomaj --bank alu --status
+-----+-----+-----+-----+
| Name   | Type(s)           | Last update status | Published release |
+-----+-----+-----+-----+
| alu    | nucleic_protein  | 2016-08-24 21:58:14 | 2003-11-26       |
+-----+-----+-----+-----+
↪-----+-----+-----+-----+
| Session          | Remote release | Release | Directory |
↪                  | Freeze        |         |           |
+-----+-----+-----+-----+
↪-----+-----+-----+-----+
| 2016-08-24 21:58:14 | 2003-11-26    | 2003-11-26 | /var/lib/biomaj/data/ncbi/
↪blast/alu/alu-2003-11-26 | no            |         |
+-----+-----+-----+-----+
↪-----+-----+-----+-----+
```

## 1.3 Advanced Configuration

Once you have this sort of simple configuration working, you may wish to explore more advanced configurations. There is a [public repository](#) of BioMAJ configurations which will be interesting to the advanced user wishing to learn more about what can be done with BioMAJ.



### 2.1 LDAP

The [BioMAJ watcher](#), provides an optional web interface to manage banks. Users can create “private” banks and manage them via the web.

### 2.2 ElasticSearch

In order to use the `--search` flag, you may wish to connect an ElasticSearch cluster.

You will need to edit your `global.properties` to indicate where the ES servers are:

```
use_elastic=0
#Comma separated list of elasticsearch nodes host1,host2:port2
elastic_nodes=localhost
elastic_index=biomaj
# Calculate data.dir size stats
data.stats=1
```

An example `docker-compose.yml` would use this:

```
version: '2'
services:
  biomaj:
    image: osallou/biomaj-docker
    links:
      - mongodb:biomaj-mongodb
      - elasticsearch
    volumes:
      - ./data:/var/lib/biomaj
      - ./global.advanced.properties:/etc/biomaj/global.properties
```

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```

mongodb:
  image: mongo

elasticsearch:
  image: elasticsearch:1.7

```

And a modified `global.properties` referenced in that file would enable elasticsearch:

```

[GENERAL]
root.dir=/var/lib/biomaj
conf.dir=%(root.dir)s/conf
log.dir=%(root.dir)s/log
process.dir=%(root.dir)s/process
cache.dir=%(root.dir)s/cache
lock.dir=%(root.dir)s/lock
#The root directory where all databases are stored.
#If your data is not stored under one directory hirearchy
#you can override this value in the database properties file.
data.dir=%(root.dir)s/data

db.url=mongodb://biomaj-mongodb:27017
db.name=biomaj

use_ldap=0
ldap.host=localhost
ldap.port=389
ldap.dn=nodomain

use_elastic=1
#Comma separated list of elasticsearch nodes host1,host2:port2
elastic_nodes=elasticsearch
elastic_index=biomaj
# Calculate data.dir size stats
data.stats=1

celery.queue=biomaj
celery.broker=mongodb://biomaj-mongodb:27017/biomaj_celery

auto_publish=1

#####
# Global properties file

#To override these settings for a specific database go to its
#properties file and uncomment or add the specific line you want
#to override.

#-----
# Mail Configuration
#-----
#Uncomment thes lines if you want receive mail when the workflow is finished

mail.smtp.host=
#mail.stmp.port=25
mail.admin=

```

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```

mail.from=biomaj@localhost
mail.user=
mail.password=
mail.tls=
# tail last X bytes of log in mail body , 0 = no tail
# mail.body.tail=2000000
# attach log file if size < X bytes, 0 for no attach
#mail.body.attach=4000000
# path to jinja template for subject, leave empty for defaults
#mail.template.subject=
# path to jinja template for body, leave empty for default
#mail.template.body=

#-----
#Proxy authentication
#-----
#proxyHost=
#proxyPort=
#proxyUser=
#proxyPassword=

#-----
# PROTOCOL
#-----
#possible values : ftp, http, rsync, local
port=21
username=anonymous
password=anonymous@nowhere.com

#access user for production directories
production.directory.chmod=775

#Number of thread during the download
bank.num.threads=4

#Number of threads to use for downloading and processing
files.num.threads=4

#to keep more than one release increase this value
keep.old.version=0

#Link copy property
do.link.copy=true

#The historic log file is generated in log/
#define level information for output : DEBUG, INFO, WARN, ERR
historic.logfile.level=INFO

http.parse.dir.line=<a[\s]+href=\"([\\S]+)/\".*alt=\"\\[DIR\\]\">.*([\\d]{2}-[\\w\\d]{2,5}-[\\d]{4})\\s[\\d]{2}: [\\d]{2})
http.parse.file.line=<a[\s]+href=\"([\\S]+)\".*([\\d]{2}-[\\w\\d]{2,5}-[\\d]{4})
\\s[\\d]{2}: [\\d]{2}) [\\s]+([\\d\\.]+)[MKG]{0,1})

http.group.dir.name=1

```

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```
http.group.dir.date=2
http.group.file.name=1
http.group.file.date=2
http.group.file.size=3

#Needed if data sources are contains in an archive
log.files=true

local.files.excluded=\\.panfs.*

#~40mn
ftp.timeout=2000000
ftp.automatic.reconnect=5
ftp.active.mode=false

# Bank default access
visibility.default=public

#proxy=http://localhost:3128

[loggers]
keys = root, biomaj

[handlers]
keys = console

[formatters]
keys = generic

[logger_root]
level = INFO
handlers = console

[logger_biomaj]
level = INFO
handlers = console
qualname = biomaj
propagate=0

[handler_console]
class = StreamHandler
args = (sys.stderr,)
level = DEBUG
formatter = generic

[formatter_generic]
format = %(asctime)s %(levelname)-5.5s [% (name)s] [% (threadName)s] %(message)s
```

API Documentation:

### 3.1 bank API reference





## 4.1 Options API reference

**class** `biomaj.options.Options` (*options=None*)  
Available options

**\_\_init\_\_** (*options=None*)  
x.**\_\_init\_\_**(...) initializes x; see `help(type(x))` for signature

**\_\_weakref\_\_**  
list of weak references to the object (if defined)

**get\_option** (*option*)  
Gets an option if present, else return None



## 5.1 Session API reference



## 6.1 Workflows API reference



## 7.1 Notify API reference





## 8.1 MetaProcess API reference

```
class biomaj.process.metaprocess.MetaProcess (bank, metas,
                                             meta_status=None,
                                             meta_data=None, simu-
                                             late=False)
```

Meta process in biomaj process workflow. Meta processes are executed in parallel.

Each meta process defined a list of Process to execute sequentially

```
__init__ (bank, metas, meta_status=None, meta_data=None, simulate=False)
```

Creates a meta process thread

**Parameters**

- **bank** (*biomak.bank*) – Bank
- **meta** (*list of str*) – list of meta processes to execute in thread
- **meta\_status** (*bool*) – initial status of the meta processes
- **simulate** (*bool*) – does not execute process

```
__get_metata_from_outputfile (proc)
```

Extract metadata given by process on stdout. Store metadata in self.metadata

**Parameters** *proc* – process

```
run ()
```

Method representing the thread's activity.

You may override this method in a subclass. The standard run() method invokes the callable object passed to the object's constructor as the target argument, if any, with sequential and keyword arguments taken from the args and kwargs arguments, respectively.

```
set_progress (name, status=None)
```

Update progress on execution

**Parameters**

- **name** (*str*) – name of process
- **status** (*bool or None*) – status of process



## 9.1 ProcessFactory API reference

```
class biomaj.process.processfactory.PostProcessFactory (bank,
                                                    blocks=None,
                                                    re-
                                                    dis_client=None,
                                                    re-
                                                    dis_prefix=None)
```

Manage postprocesses

self.blocks: dict of meta processes status Each meta process status is a dict of process status

```
__init__ (bank, blocks=None, redis_client=None, redis_prefix=None)
```

Creates a postprocess factory

**Parameters**

- **bank** (biomaj.bank.Bank) – Bank
- **blocks** (*dict*) – initial status of block processes

```
run (simulate=False)
```

Run processes

**Parameters** **simulate** (*bool*) – does not execute process

**Returns** status of execution - bool

```
class biomaj.process.processfactory.PreProcessFactory (bank,
                                                    metas=None, re-
                                                    dis_client=None,
                                                    re-
                                                    dis_prefix=None)
```

Manage preprocesses

```
__init__ (bank, metas=None, redis_client=None, redis_prefix=None)
```

Creates a preprocess factory

**Parameters**

- **bank** (biomaj.bank.Bank) – Bank

- **metas** (*dict*) – initial status of meta processes

**run** (*simulate=False*)

Run processes

**Parameters** **simulate** (*bool*) – does not execute process

**Returns** status of execution - bool

**class** `biomaj.process.processfactory.ProcessFactory` (*bank*, *re-*  
*dis\_client=None*,  
*redis\_prefix=None*)

Manage process execution

**\_\_init\_\_** (*bank*, *redis\_client=None*, *redis\_prefix=None*)

`x.__init__(...)` initializes `x`; see `help(type(x))` for signature

**\_\_weakref\_\_**

list of weak references to the object (if defined)

**fill\_tasks\_in\_threads** (*metas*)

Dispatch meta processes in available threads

**run** (*simulate=False*)

Run processes

**Parameters** **simulate** (*bool*) – does not execute process

**Returns** status of execution - bool

**run\_threads** (*simulate=False*)

Start meta threads

**Parameters** **simulate** (*bool*) – do not execute processes

**Returns** tuple global execution status and status per meta process

**class** `biomaj.process.processfactory.RemoveProcessFactory` (*bank*,  
*metas=None*,  
*re-*  
*dis\_client=None*,  
*re-*  
*dis\_prefix=None*)

Manage remove processes

**\_\_init\_\_** (*bank*, *metas=None*, *redis\_client=None*, *redis\_prefix=None*)

Creates a remove process factory

**Parameters**

- **bank** (`biomaj.bank.Bank`) – Bank
- **metas** (*dict*) – initial status of meta processes

**run** (*simulate=False*)

Run processes

**Parameters** **simulate** (*bool*) – does not execute process

**Returns** status of execution - bool

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