# nebulizer Documentation

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# Nebulizer is a Python utility which provides a high-level interactive command-line interface to remotely administer Galaxy servers.

Nebulizer was developed as a tool to help "part-time" Galaxy admins perform day-to-day administrative tasks across multiple Galaxy instances, enabling various admin operations to be executed via the command line as an alternative to using the Galaxy web interface.

Nebulizer is built on top of the Bioblend library and offers an interface and range of functionality that complements the lower-level interfaces offered by Ephemeris and Parsec.

Contents:

# nebulizer

Command-line utilities to help with managing users, data libraries and tools in a Galaxy instance, using the Galaxy API via the Bioblend library.

- Free software: Academic Free License version 3.0
- Documentation: https://nebulizer.readthedocs.io
- Code: https://github.com/pjbriggs/nebulizer

#### Note: Nebulizer is still a work in progress.

Please exercise caution when attempting irreversible operations, especially against production Galaxy instances (for example when creating users or data libraries).

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**Quick Start** 

This quick start gives some examples of using nebulizer commands to perform remote administration tasks on a Galaxy instance from the command line.

## 1.1 Getting Nebulizer

It is recommended to install Nebulizer via pip in a virtualeny, for example:

```
% virtualenv .venv
% source .venv/bin/activate
% pip install nebulizer
```

This will provide an executable called nebulizer with a number of subcommands for performing different tasks remotely on Galaxy instances.

#### 1.2 Nebulizer Basics

Generally Nebulizer commands take the form:

```
nebulizer COMMAND GALAXY [OPTIONS]
```

To interact remotely with a Galaxy instance using Nebulizer requires at minimum the URL of the instance and then either an API key or a user login name.

For example to list the data libraries available on Galaxy Main:

```
nebulizer -k 9b376af2250818d14949b3c list_libraries https://usegalaxy.org
```

or

```
nebulizer -u USER@DOMAIN list_libraries https://usegalaxy.org
```

In this second case Nebulizer will prompt for the Galaxy password to authenticate the user login, unless it's supplied via the -P option.

To store the Galaxy URL-API key pair against an alias main, to avoid needing full authentication details each time:

```
nebulizer add_key main https://usegalaxy.org 9b376af2250818d14949b3c
```

or alternatively get Nebulizer to fetch the API key itself by supplying the user login:

```
nebulizer -u USER@DOMAIN add_key main https://usegalaxy.org
```

More information on managing API keys in Nebulizer can found here.

The stored alias is then used in subsequent commands, for example to list the data libraries again it is now sufficient to do just:

```
nebulizer list_libraries main
```

The following sections contain examples of how Nebulizer might be used to perform various administrive tasks.

Nebulizer provides subcommands to perform various administrive tasks:

#### Managing users:

- list\_users
- create\_user
- create\_batch\_users
- create\_users\_from\_file
- delete\_user

#### Managing data libraries:

- list\_libraries
- create\_library
- create\_library\_folder
- add\_library\_datasets

#### Managing tools:

- list\_tools
- list\_tool\_panel
- install\_tool
- update\_tool
- uninstall\_tool
- search\_toolshed

#### Managing quotas:

- quotas
- quota\_add
- quota\_mod

• quota\_del

Querying Galaxy instances:

- ping (check if a Galaxy instance is alive)
- config (fetch configuration for a Galaxy instance)

See the tutorial for a walkthrough some of these commands.

#### 1.3 Contributors

Nebulizer has been developed by Peter Briggs @pjbriggs, with contributions from:

• Hugo van Kemenade (@hugovk)

Thanks to Peter van Heuseden (@pvanheus) for porting Nebulizer into Bioconda.

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Installation

## 2.1 Virtualenv and pip

For a traditional installation of Nebulizer, first set up a Python virtualenv (this example creates a new one in .venv) and then install with pip:

```
$ virtualenv .venv;
$ source .venv/bin/activate
$ pip install nebulizer
```

When installed this way, Nebulizer can be upgraded as follows:

```
$ . .venv/bin/activate
$ pip install -U nebulizer
```

To install or update to the latest development branch of Nebulizer with pip, use the following pip install idiom instead:

```
$ pip install -U git+git://github.com/pjbriggs/nebulizer.git@devel
```

Nebulizer should work with recent Python 3 versions.

#### 2.2 Conda

Nebulizer can be installed using Conda (most easily obtained via the Miniconda Python distribution):

```
$ conda config --add channels bioconda
$ conda install nebulizer
```

Note that the version available via bioconda may lag the most recent version available via pip.

#### Configuration

Nebulizer doesn't require any additional configuration after installation: it is possible to supply either a Galaxy API key or a Galaxy username/password combination on the command line (alongside the URL of the Galaxy instance) each time a command is executed.

However this is both laborious and potentially insecure. Nebulizer can store Galaxy URL-API key pairs against aliases locally to make this easier, using the add\_key command.

The simplest way to create a new alias is:

```
nebulizer -u EMAIL add_key ALIAS GALAXY_URL
```

Nebulizer will prompt for the password for the account and will then fetch and store the API key automatically.

**Note:** For example: to add an alias main for Galaxy main at https://usegalaxy:

```
nebulizer -u peter.briggs@manchester.ac.uk add_key main https://usegalaxy.org
```

Alternatively you can supply an API directly on the command line:

```
nebulizer add_key ALIAS GALAXY_URL API_KEY
```

#### **Note:** For example:

```
nebulizer add_key main https://usegalaxy.org 9b376af2250818d14949b3c
```

The stored alias can then be used instead of supplying the full Galaxy URL with an email/password combination or an API key; for example to get configuration information from Galaxy main using the config command:

```
nebulizer config main
```

See Storing and managing Galaxy API keys for more information on managing stored aliases.

Managing Users

## 4.1 Querying user information

list\_users displays user emails and names in a Galaxy instance:

nebulizer list\_users GALAXY

- -1: returns extended information for each user (status, whether they are an admin user, disk usage and quota size and usage).
- --status: filter list on user status, which can be one of active (the default), deleted (only list deleted users which haven't been purged) or all (list all active, deleted, and purged users).
- --sort: specify one or more fields to sort the output on; valid fields are email, disk\_usage, quota, quota\_usage. Multiple fields should be separated by commas (e.g. --sort=quota, disk\_usage).
- --name: filter list on user email (can include glob-style wildcards e.g. --name="\*blogqs\*").

## 4.2 Creating and deleting individual users

create\_user makes a new user account:

```
nebulizer create_user GALAXY USER@DOMAIN [ PUBLIC_NAME ]
```

If PUBLIC\_NAME is not supplied then one will be generated automatically from the email address.

- -c: check if the account already exists
- -p: supply password for the new account (otherwise Nebulizer will prompt for a password)

delete\_user removes a user account:

```
nebulizer delete_user GALAXY USER@DOMAIN
```

• --purge: purge the account as well as deleting (or purge an unpurged account which has been previously deleted)

**Note:** Purging a user account overwrites the email and username for that account with random strings; it also marks the datasets and histories associated with that account as deleted. These data can then be removed from disk by running Galaxy's clean-up scripts.

For information on the clean-up scripts see the Galaxy documentation at https://galaxyproject.org/admin/config/performance/purge-histories-and-datasets/

#### 4.3 Creating batches of users from a template name

create\_batch\_users makes a set of user accounts based on a template name:

```
nebulizer create_batch_users GALAXY TEMPLATE [START] END
```

The template email address should include a # symbol which acts as as a placeholder for an integer index, e.g.

```
user#@example.org
```

The integer indices are generate from the range START...END (if START is not supplied then it is set to 1).

For example:

```
nebulizer create_batch_users user#@example.org 1 5
```

#### creates accounts:

```
user1@galaxy.org
user3@galaxy.org
user4@galaxy.org
user5@galaxy.org
```

- -c: check if the accounts already exist
- -p: supply password for the new accounts (otherwise Nebulizer will prompt for a password); the same password will be applied to all the new accounts

#### 4.4 Creating user accounts from a file

create\_users\_from\_file makes a set of user accounts using data from a file:

```
nebulizer create_users_from_file GALAXY USERS_FILE
```

USER\_FILE is a tab-delimited file with the following fields on each line:

```
email|password|public_name
```

defining a new account.

**Note:** Optionally public\_name can be left out and will then be generated automatically.

• -c: check if the accounts already exist

#### Managing Data Libraries

## 5.1 Querying data libraries

list\_libraries displays information about the data libraries and their contents in a Galaxy instance:

```
nebulizer list_libraries GALAXY [ LIBRARY[/PATH...] ]
```

By default all the libraries in the instance are listed; if the name of a data library (LIBRARY) is included then the datasets and folders in that library are listed. If a PATH (names of subfolders separated by /) is supplied then the contents of the matching folder are displayed.

For example: listing the data libraries in Galaxy main:

```
nebulizer list_libraries https://usegalaxy.org
```

List the contents of the "Tutorials" data library:

```
nebulizer list_libraries https://usegalaxy.org Tutorials
```

List the contents of the "ChIP\_seq (Reb1)" folder in the "Tutorials" data library:

```
nebulizer list_libraries https://usegalaxy.org "Tutorial/ChIP_seq (Reb1)"
```

• -1: list extended information about libraries, folders and datasets

## 5.2 Creating and populating data libraries

create\_library creates a new data library:

```
nebulizer create_library GALAXY LIBRARY_NAME [--description DESCRIPTION] [--synopsis_ + SYNOPSIS]
```

create\_library\_folder creates a new folder in an existing data library or folder:

```
nebulizer create_library_folder GALAXY LIBRARY_NAME/[PATH]
```

For example: create a data library called NGS data and a folder Run 21 within it:

```
nebulizer create_library GALAXY "NGS data 2020" \
--description="Sequencing data analysed in 2020"
nebulizer create_library_folder GALAXY "NGS data 2020/Run 21"
```

add\_library\_datasets uploads datasets to a data library:

```
nebulizer add_library_datasets GALAXY LIBRARY_NAME/PATH FILE [FILE...]
```

- --file-type: specify the Galaxy data type to assign to datasets
- --dbkey: specify the dbkey for the datasets
- --server: upload files from the filesystem of the server that Galaxy is running on (default is to upload from the local filesystem)
- --link: create symlinks to the files on the server (if --server is also specified)

For example, add Fastq files to a data library folder:

```
nebulizer add_library_datasets galaxy "NGS data/Run 21" ~/Sample1_R*.fq \
--file-type=fastqsanger --dbkey=hg38
```

**Managing Tools** 

## 6.1 Querying tool information

By default, list\_tools displays information on all the tool repositories that are installed in a Galaxy instance:

```
nebulizer list_tools GALAXY
```

For each installed repository the details include: repository name, toolshed, owner, revision id and changeset, and installation status.

Repository details are also preceded by a single-character 'status' indicator:

- D = deprecated;
- ^ = newer revision is also installed;
- u = update (newer revision) is available but not installed;
- U = upgrade available (new revision with a change of tool version) but not installed;
- \* = latest revision installed

list\_tools supports a number of options to modify its behaviour, including:

- --updateable: only list tool repositories that have uninstalled available updates or upgrades
- --built-ins: include details of the "built-in" tools within the Galaxy instance (i.e. those not installed from a toolshed)

An alternative 'tool-centric' view of the tools in a Galaxy instance can be obtained using the --mode=tools option.

**Note:** This is a new version of the list\_tools which replaces the old list\_installed\_tools command (which is no longer available). The --mode=tools option replicates the output from the old list\_tools command.

list\_tool\_panel displays information on the tool panel sections in a Galaxy instance:

```
nebulizer list_tool_panel GALAXY
```

New tool panel sections can be created when installing tool repositories.

#### 6.2 Install, update and remove tools

install\_tool installs tools from a Galaxy toolshed (by default the main Galaxy toolshed at https://toolshed.g2.bx.psu.edu):

```
nebulizer install_tool GALAXY TOOL_REPOSITORY
```

The tool repository can be specified as:

- [TOOLSHED] OWNER/TOOLNAME [ REVISION ] e.g. devteam/fastqc (defaults to most recent revision from the main toolshed), or devteam/fastqc e7b2202befea (specifies revision e7b2202befea)
- toolshed URL with or without a revision (e.g. https://toolshed.g2.bx.psu.edu/view/devteam/fastqc/e7b2202befea)
- --tool-panel-section: specify a tool panel section to install the tools from the repository into (otherwise tools appear at the "top level" of the tool panel)

For example: to install the most recent FastQC tool from the main Galaxy toolshed under the NGS: QC and manipulation section of the tool panel:

```
nebulizer install_tool GALAXY devteam/fastqc \
    --tool-panel-section="NGS: QC and manipulation"
```

To install a specific revision of the Trimmer tool from the test toolshed:

```
nebulizer install_tool GALAXY \
  https://testtoolshed.g2.bx.psu.edu/view/devteam/trimmer/dec27ea206c3 \
  --tool-panel-section="Test tools"
```

update\_tool installs the latest revision of a previously installed tool repository, if a new version is available:

```
nebulizer update_tool GALAXY TOOL_REPOSITORY
```

The tool repository is specified as with install\_tool except that a revision cannot be included. For example:

```
nebulizer update_tool GALAXY devteam/fastqc
```

It is also possible to include glob-style wildcards in the tool repository name and/or owner e.g. devteam/\* or  $bgruening/deeptools_*$ . To request update of all tools:

```
nebulizer update_tool GALAXY '*/*'
```

Note: update\_tool doesn't uninstall the older versions of the tools that are updated.

**Warning:** By default checks on the availability of updates for tools performed by the list\_tools and update\_tool commands are done using information cached by the Galaxy instance in question. As a result these commands may not always indicate when updates are available.

To force these commands to check the installed revisions against those in the toolshed, add the --check-toolshed option. Note however that this can impose a significant overhead which can make the commands much slower.

uninstall\_tool removes a previously installed tool:

```
nebulizer uninstall_tool GALAXY TOOL_REPOSITORY
```

The tool repository is specified as with install\_tool, for example to uninstall and deactivate a specific revision of a tool:

```
nebulizer uninstall_tool GALAXY devteam/fastqc/e7b2202befea
```

To uninstall all installed revisions of a tool and remove from disk:

```
nebulizer uninstall_tool localhost devteam/fastqc/* \
   --remove_from_disk
```

## 6.3 Searching for tool repositories on a Toolshed

search\_toolshed searches for tools on a toolshed:

```
nebulizer search_toolshed QUERY
```

QUERY can include glob-style wildcards. For example, to search the main toolshed for Deeptools related tools:

```
nebulizer search_toolshed "deeptools_*"
```

• --toolshed: specify the URL of the toolshed to search.

## 6.4 Bulk tool repository management

install\_tool --file installs the tool repositories listed in a tab-delimited file into a Galaxy instance:

```
nebulizer install_tool GALAXY TOOLS_FILE
```

TOOLS\_FILE must be a tab-delimited list of repositories, one repository per line in the format:

```
TOOLSHED | OWNER | REPOSITORY | REVISON | SECTION
```

#### For example:

```
toolshed.g2.bx.psu.edu devteam bowtie_wrappers 9ca609a2a421 NGS: Mapping
```

list\_tools --mode=export can generate a list of tool repositories already installed in a Galaxy instance in this format, e.g.:

```
nebulizer list_tools GALAXY --mode=export
```

By combining these two commands it is possible to 'clone' the installed tools from one Galaxy instance into another.

For example to replicate the tools installed on the 'Palfinder' instance into a local Galaxy:

```
nebulizer list_tools https://palfinder.ls.manchester.ac.uk --mode=export > palfinder.

→tsv

nebulizer install_tool http://127.0.0.1 --file palfinder.tsv
```

**Warning:** Bulk installation of tools in this manner should be used with caution, especially when installing into a Galaxy instance which already has installed tools.

#### Managing Quotas

## 7.1 Querying quotas

quotas lists the quotas defined in a Galaxy instance:

nebulizer quotas GALAXY

- -1: returns extended information for each quota, including the associated users and groups.
- --status: filter list on quota status, which can be one of active (the default), deleted (only list deleted quotas) or all (list all active and deleted quotas).
- --name: filter list on quota name (can include glob-style wildcards e.g. --name="\*NGS\*").

## 7.2 Creating and deleting quotas

quota\_add defines a new quota:

```
nebulizer quota_add GALAXY QUOTA_NAME SIZE
```

SIZE can either be an amount (e.g. 10GB, 0.2 T) or an amount preceded by an operation (one of +, - or =, e.g. = 300gb, +100G). If an operation isn't specified then = is assumed.

- -d/--description: set the description for the quota (defaults to the quota name).
- --default\_for: set the quota as the the default for either 'registered' or 'unregistered' users.

Users and groups can be associated with the new quota using the -u and -q options:

- -u/--users: associate one or more users with the quota, as a comma-separated list of email addresses.
- -g/--groups: associate one or more groups with the quota, as a comma-separated list of group names.

quota\_del deletes an existing quota:

nebulizer quota\_del GALAXY QUOTA\_NAME

Note: A deleted quota can be restored using the --undelete option of the quota\_mod command.

## 7.3 Modifying quota definitions

quota\_mod updates a quota definition:

```
nebulizer quota_mod GALAXY QUOTA_NAME ...
```

Options allow various quota properties to be modified:

- -n/--name: sets a new name for the quota.
- -d/--description: sets a new description.
- -q/--quota-size: updates the size of the quota, and how it is applied.
- --default\_for: set the quota as the the default for either 'registered' or 'unregistered' users.

Users and groups can be associated with the following options:

- -a/--add-users: associate one or more users with the quota, as a comma-separated list of email addresses.
- -r/--remove-users: disassociate one or more users from the quota, as a comma-separated list of email addresses.
- -M--add-groups: associate one or more groups with the quota, as a comma-separated list of group names.
- -R/--remove-groups: disassociate one or more groups from the quota, as a comma-separated list of group names.

Previously deleted quotas can be restored using the -u/--undelete option.

Querying Galaxy instances

## 8.1 Checking status and configuration of a Galaxy server

'Ping' a Galaxy instance to check it's alive and responding to requests:

nebulizer ping GALAXY

Get information about an instance's configuration using

:: nebulizer config GALAXY

## Storing and managing Galaxy API keys

The majority of Nebulizer's commands require valid credentials in order to interact with a Galaxy instance. These can be supplied explicitly on the command line each time:

```
nebulizer -u USER@DOMAIN [ -P PASSWORD ] ...command...
nebulizer -k API_KEY ...command...
```

Alternatively it is possible to store Galaxy URL-API key pairs in a file called .nebulizer located in the user's home directory, with each pair being associated with an alias.

The file is created automatically the first time an alias is created using add\_key command, and consists of tabdelimited lines with three fields:

```
alias|Galaxy_URL|API_key
```

#### for example:

```
demo http://127.0.0.1:8080 4551fbf7cd8b1bc59db....
```

This file can be manually edited using a text editor such as vi; however Nebulizer also provides a set of commands for querying and modifying the file contents.

list\_keys shows the aliases with their associated Galaxy URLs:

```
nebulizer list_keys
```

Note: By default the API keys are not shown by list\_keys; use the --show-api-keys option to include them.

**Note:** Use the whomai command to find out which user is associated with an alias:

```
nebulizer whoami ALIAS
```

add\_key will store a Galaxy-API key combination under a new alias. If the API key is known then the general form of the command is:

```
nebulizer add_key ALIAS GALAXY_URL API_KEY
```

However it is usually easier to get Nebulizer to fetch the key automatically, by supplying a Galaxy username (email):

```
nebulizer -u USER@DOMAIN add_key ALIAS GALAXY_URL
```

Multiple Galaxy URL-key pairs can be stored; only the associated aliases need to be unique.

update\_key will update the details stored for an existing alias:

```
nebulizer update_key ALIAS --new-url GALAXY_URL
nebulizer update_key ALIAS --new-api-key API_KEY
nebulizer -u USER@DOMAIN update_key ALIAS --fetch-api-key
```

remove\_key deletes an existing ALIAS and associated credentials:

```
nebulizer remove_key ALIAS
```

## General options

Nebulizer supports a number of general options for managing how connections are made to Galaxy servers, how to authenticate, and what information to output.

#### 10.1 Authenticating without stored credentials

Use --api\_key (-k) to explicitly specify the API key to use for authentication on the command line, e.g.

```
nebulizer -k API_KEY COMMAND...
```

Use the --username (-u) option to authenticate using the normal Galaxy login credentials (i.e. email and password) instead of the API key, e.g.

```
nebulizer -u USER@DOMAIN COMMAND...
```

You will be prompted to enter the password, but you can also use the  $--galaxy\_password(-P)$  option to specify it explicitly on the command line.

## 10.2 Controlling warnings and debugging output

--suppress-warnings (-q) suppresses warning messages from Nebulizer; conversely debugging output can be enabled using the --debug option.

## 10.3 Handling SSL certificate verification failures

Nebulizer commands will fail for Galaxy instances which are served over https protocol without a valid SSL certificate, reporting an error like:

```
[SSL: CERTIFICATE_VERIFY_FAILED] certificate verify failed (_ssl.c:590), 0 attempts_

→left: None
```

In this case adding the --no-verify(-n) option turns off the certificate verification and should enable a connection to be made.

**Nebulizer Tutorial** 

This tutorial offers a short hands-on introduction to the main functionality offered by Nebulizer.

## 11.1 0. Preparation: making a temporary local Galaxy instance

Before starting it is recommended to set up a temporary local Galaxy instance to run the tutorial on, rather than trying Nebulizer out on an existing installation.

The following commands should fetch and start up the local instance:

```
# Get the Galaxy source and put into a new directory
# called 'galaxy_for_demo'
git clone -b release_21.01 https://github.com/galaxyproject/galaxy.git galaxy_for_demo
# Move into the source directory
cd galaxy_for_demo
# Make a configuration file
cp config/galaxy.yml.sample config/galaxy.yml
```

You will need to edit the configuration file <code>config/galaxy.yml</code> using a text editor (e.g. vi, nano, <code>gedit</code> etc) to specify an admin user, by changing the line:

```
#admin_users: null
```

to e.g.

```
admin_users: admin@localhost.org
```

or whatever email address you want. You will also need to change the line:

```
#allow_user_deletion: false
```

to

```
allow_user_deletion: true
```

Then start Galaxy running:

```
sh run.sh
```

**Note:** This step can take some time when it's first run; however once Galaxy has installed its dependencies and performed its initial configuration then subsequent startups of the demo Galaxy should be much quicker.

Use ctrl-C to stop Galaxy running, and repeat this command to start it up again.

Once Galaxy is running you can connect to it by pointing a web browser to http://127.0.0.1:8080/

Finally you will need to register an account in Galaxy with the same email address as the one used for admin\_users (admin@localhost.org in this example), using the web browser.

Now you're ready to run through the tutorial.

**Note:** Full instructions for setting up a local Galaxy instance can be found at https://galaxyproject.org/admin/get-galaxy/

#### 11.2 1. Install Nebulizer

Installing Nebulizer is best done in a Python virtual environment - for example:

```
virtualenv -p python3 venv.nebulizer
. venv.nebulizer/bin/activate
pip install nebulizer
```

Note: It is recommended to do this in a different directory to the local Galaxy instance from the previous section.

Once this is done you should have access to the nebulizer utility.

To list the available commands:

```
nebulizer --help
```

You use the ping command to check if a Galaxy instance is active. For example to check the main Galaxy server:

```
nebulizer ping https://usegalaxy.org
```

which should produce output like:

```
https://usegalaxy.org: status = ok time = 574.041 (ms)
https://usegalaxy.org: status = ok time = 587.883 (ms)
https://usegalaxy.org: status = ok time = 549.526 (ms)
...
```

**Note:** Do control-C to terminate the "ping".

You can also use the config command to query the details of a Galaxy instance's configuration. For example to query the local Galaxy:

nebulizer config http://127.0.0.1:8080

#### 11.3 2. Set up aliases for Galaxy API keys

Most Nebulizer commands require you to interact with a Galaxy instance using an account on that instance.

For these commands you can authorise access each time by specifying your registered email address and password or Galaxy API key on the command line. For example:

nebulizer -u admin@localhost.org whoami http://127.0.0.1:8080

**Note:** -u will prompt you to enter the password for the account before performing the action; -k can be used to specify the API key.

Warning: This won't work if you didn't make an account for admin@localhost.org when preparing the local Galaxy in the previous step!

This is quite laborious when executing several commands, so Nebulizer allows you to associate Galaxy instances and their API keys with aliases; these are used as shortcuts when running the commands.

To see the aliases and associated Galaxy servers:

nebulizer list\_keys

**Note:** If you've never used Nebulizer before then nothing will be listed.

To set up a new alias called local and associate it with the admin account in our local Galaxy, we can do:

nebulizer -u admin@localhost.org add\_key local http://127.0.0.1:8080

This will prompt you for the password for the account and then create the alias. Once this is done you can repeat the list\_keys command and see an entry for the local Galaxy:

local http://127.0.0.1:8080

In subsequent commands you can use local rather than specifying the full Galaxy URL, and won't need to enter your email or password. For example:

nebulizer whoami local

Now we're ready to do some basic administration of our local Galaxy using Nebulizer.

Note: See Storing and managing Galaxy API keys for more details.

#### 11.4 3. Listing, adding and deleting users

We can list the users in our local Galaxy with:

```
nebulizer list_users local
```

There will be just one account (the original admin account we made at the start).

We can add a new user using:

```
nebulizer create_user local ann.onymous@manchester.ac.uk
```

**Note:** This will prompt you for a password for the new account; use the -p option to set the password via the command line.

Do the list\_users command again to see new user listed. Use the -1 option to display additional information about each user is displayed, including status and disk usage (and quota usage, if quotas are enabled).

Batches of user accounts can be created from a "template" name using the create\_batch\_users command; this can be useful for example when setting up Galaxy instances for teaching:

```
nebulizer create_batch_users local user#@bcc2020.org 5
```

**Note:** This will prompt you for a password which will be assigned to all the new accounts.

Use the list\_users command to see the new accounts:

```
      user1@bcc2020.org
      user1

      user2@bcc2020.org
      user2

      ...
      user5@bcc2020.org
```

Accounts can also be deleted:

```
nebulizer delete_user local user5@bcc2020.org
```

The user will no longer be listed by list\_users.

**Warning:** If the deletion fails then check that the Galaxy configuration has allow\_user\_deletion set to true.

**Note:** See *Managing Users* for more details.

## 11.5 4. Creating and populating data libraries

We can list the data libraries in our local Galaxy instance using:

nebulizer list\_libraries local

Initially our local Galaxy doesn't contain any library data; we can create a new data library using:

nebulizer create\_library local "Example data"

Note: Use the -d and -s options to add description and synopsis information for the new library.

Now this will be listed by the list\_libraries command. We can list the contents of a library by specifying its name:

nebulizer list\_libraries local "Example data"

Initially the library is empty; we can create a folder within the library:

nebulizer create\_library\_folder local "Example data/Fastqs"

To list the contents of a library folder specify the "path" to the folder:

nebulizer list\_libraries local "Example data/Fastqs"

Datasets can be added to libraries and folders from the local workstation:

 $\verb|nebulizer| add_library_datasets| local "Example data/Fastqs" Illumina_SG_R* --dbkey=hg38| local "Example data/Fastqs" --dbkey=hg38| local "Example data/Fastqs$ 

**Note:** The example Fastq files can be found here:

- Illumina\_SG\_R1.fastq
- Illumina\_SG\_R2.fastq

When listing the contents of libraries and folders, additional information is reported by specifying the -1 option:

nebulizer list\_libraries local "Example data/Fastqs" -1

**Note:** See *Managing Data Libraries* for more details.

## 11.6 5. Installing and managing tools

We can list the tools installed in our local Galaxy using:

nebulizer list\_tools local

Initially there are no tools installed; we can search the main Galaxy toolshed for the tools we want to install, for example the FastQC tool:

nebulizer search\_toolshed fastqc

**Warning:** The time taken for searching depends on the speed of the toolshed, so sometimes this can be slow if e.g. the toolshed is experiencing issues.

This will list all the tool repositories and toolshed versions available to install:

```
devteam fastqc 21:e7b2202befea devteam fastqc 19:9da02be9c6cc devteam fastqc 16:ff9530579d1f ...
```

We can install the latest version of FastQC with

```
nebulizer install_tool local devteam/fastqc --tool-panel-section="NGS tools"
```

Note: Using --tool-panel-section will create a new section in the Galaxy tool panel and put the tools from this repository under it; otherwise tools are not installed under any section. You can use the list\_tool\_panel command to see what tool panel sections are already present.

Running list\_tools now shows the tool repository is installed:

```
* fastqc toolshed.g2.bx.psu.edu devteam 21:e7b2202befea Installed
```

**Note:** The \* next to tool repository indicates that this is most recent version.

Use the --mode=tools option to list the associated tools instead.

We can install a specific version of a tool repository, for example the Trimmomatic tool:

```
nebulizer install_tool local pjbriggs/trimmomatic 51b771646466 --tool-panel-section=
→"NGS tools"
```

Running list\_tools now shows this tool repository is also installed:

```
* fastqc toolshed.g2.bx.psu.edu devteam 21:e7b2202befea Installed U trimmomatic toolshed.g2.bx.psu.edu pjbriggs 12:51b771646466 Installed
```

Here U indicates there is a newer revision available with a new version of the tool (u indicates a newer revision without a tool version update).

Rerunning the list\_tools command with the --updateable option filters the list of tool repositories to just those with available updates.

We can update Trimmomatic to the newest version automatically by running the update\_tool command:

```
nebulizer update_tool local pjbriggs/trimmomatic
```

**Note:** This installs the most recent version but doesn't remove the older version.

The uninstall\_tool command removes an installed repository; for example to uninstall the older Trimmomatic tool version:

nebulizer uninstall\_tool local pjbriggs/trimmomatic 51b771646466

Running  $\mathtt{list\_tools}$  shows that the older tool repository is no longer present.

**Note:** See *Managing Tools* for more details.

## History

#### 12.1 v0.7.1 (2021-06-29)

- Pin required bioblend version to 0.15.0 (PR #40).
- Switched CI tests to Github Actions (PR #40).

## 12.2 v0.7.0 (2021-05-17)

#### **Breaking changes:**

- Dropped support for Python 2.7: nebulizer now needs Python 3.6+ (thanks to Hugo van Kemenade @hugovk) (PR #102)
- Substantial refactoring and simplification of the tool managament commands (PR #113): Now only list\_tools, install\_tool, update\_tool

and delete\_tool commands are supported

- list\_installed\_tools renamed to list\_tools
- old functionality of list\_tools replaced by list\_tools --mode=tools
- list\_repositories command dropped; functionality replaced by list\_tools
  --mode=export
- install\_repositories command dropped; functionality replaced by install\_tool
   --file=...

#### New commands:

• New quota, quota\_add, quota\_mod and quota\_del commands for managing quotas (PR #66)

#### **Updates to existing commands:**

• -l option for search\_toolshed includes the shed URL (PR #91)

- update\_tool allows use of wildcards (i.e. \*) when specifying tool repository names and owners, to enable
  multiple tool repositories to updated at once (PR #92)
- New --status option for list\_users command allows deleted and purged user accounts to also be listed (PR #97)
- New --sort option for list\_users command allows sorting of listed accounts by disk usage, quota and quota usage (PR #104)

#### **Bug fixes:**

- Fix to using the --purge option of the delete\_user command (previously it wasn't possible to purge accounts) (PR #98)
- Remove requirement to specify an account or API key on Galaxy server when using the ping and config commands (PR #100)

#### Other updates:

• Added support for Python 3.9 (thanks to Hugo van Kemenade @hugovk) (PR #108)

#### 12.3 v0.6.0 (2020-07-14)

#### New commands:

- New search toolshed command (PR #42)
- New config command (PR #57)
- New delete\_user command (PR #62)
- New uninstall\_tool command (PR #64)

#### **Updates to existing commands:**

- list\_keys doesn't report API keys unless --show-api-keys option is specified (PR #58)
- Additional fields reported by --long-listing-format option of list\_users (disk and quota usage, status); doesn't report ID by default (PR #59)
- Enable flexible tool repository specification syntax for install\_tool and update\_tool (PR #60)
- remove\_key prompts user to confirm API key deletion (PR #72)
- Use spaces rather than tabs to line up fields in output from list\_users, list\_installed\_tools, list\_tools, list\_tool\_panel, list\_keys, config, list\_libraries; use --show\_id to report Galaxy IDs for users and data libraries (PR #68, PR #69, PR #70)

#### **Documentation:**

- Add a tutorial/walkthrough (PR #75)
- Significant overhaul and expansion of documentation (PR #78)

#### Removed functionality:

• Removed deprecated utilities manage\_users, manage\_tools and manage\_libraries (PR #61)

#### 12.4 v0.5.0 (2020-04-20)

Add support for Python 3.6, 3.7 and 3.8 (PR #50, PR #51)

#### 12.5 v0.4.3 (2018-10-05)

• Ensure that click dependency is version 6.7 or earlier, to avoid subcommand names changing from e.g. list\_users to list-users (PR #49)

#### 12.6 v0.4.2 (2017-08-24)

- Commands now explicitly return appropriate exit code values indicating success (0) or failure (non-zero values).
- New option —check—toolshed added to list\_installed\_tools and update\_tool commands, to check installed revisions directly against those available in the toolshed (PR #41)
- Update install\_tool, update\_tool and install\_repositories to install tool dependencies through a resolver (e.g. conda) by default (issue #43)
- New options added to install\_tool, update\_tool and install\_repositories commands, to explicit control how tool and repository dependencies should be handled (PR #44):
  - --install-tool-dependencies [yes|no]: install tool dependencies via the toolshed, if any are defined (default is yes)
  - --install-tool-dependencies [yes|no]: install tool dependencies via the toolshed, if any are defined (default is yes)
  - --install-resolver-dependencies [yes|no]: install dependencies through a resolver that supports installation (e.g. conda) (default is yes)

## 12.7 v0.4.1 (2016-12-19)

• Fix broken update\_tool command (PR #40).

## 12.8 v0.4.0 (2016-11-18)

- New subcommand ping: 'ping' a Galaxy instance to see if it's responsive (PR #33).
- New subcommand whoami: reports user associated with the API key (PR #37).
- add\_library\_datasets: refuses to perform upload if using the master API key (essentially API key must have an associated user).
- install\_repositories: prints a list of all tool repositories that couldn't be installed.
- New --timeout and --nowait options added for install\_tool, update\_tool and install\_repositories subcommands.
- Fix to treat tool repositories with status New as still installing when trying to install tools (PR #31).
- Some improvements to logging (PR #38).

## 12.9 v0.3.0 (2016-10-26)

• New class tools. ToolPanel and updates to existing tools. ToolPanelSection class.

- install\_tool: fix behaviour so that command does nothing if a version is not specified and at least one version of the tool is already installed.
- list\_repositories and install\_repositories: new commands to generate a list of installed tool repositories from a Galaxy instance and then reinstall tool repositories from a list with the same format (PR #19).
- install\_tool: fix incorrect reporting of target tool panel section (PR #20)
- add\_key and update\_key: fix automatic retrieval of API key, which only worked previously if connecting
  user was an admin account (PR #23)
- list\_tool\_panel: shows tools in order they appear in Galaxy when using --list-tools option.
- Deprecated utilities (manage\_users, manage\_tools and manage\_libraries) issue warnings when run.
- License updated to Academic Free License (AFL).
- Initial version of documentation also made available via ReadTheDocs (PR #21)

#### 12.10 v0.2.0 (2016-10-17)

- Implemented new nebulizer utility which provides all previous functionality via subcommands, plus commands for managing API keys automatically (old manage\_users, manage\_tools and manage\_libraries utilities are still available for backwards-compatibility but are deprecated).
- New general options:
  - -q/--suppress-warnings: prevent warning messages from nebulizer commands.
- Various fixes and improvements to underlying functionality:
  - install\_tools: now checks if tool is already installed; handles tool revisions that include the revision number; polls Galaxy until tool is installed, or operation times out; exit status reflects the success or failure of the installation.
  - update\_tool: now works even if original tool isn't in a tool panel section
  - list\_installed\_tools: now groups tools under correct repo revision when using --list-tools option.

## 12.11 v0.1.1 (2016-05-16)

• Add -u/--username and -P/--galaxy\_password options to all commands to allow interaction with Galaxy instance via API using normal login credentials instead of API key.

#### 12.12 v0.1.0 (2015-11-06)

• Initial release of nebulizer utilities for administering Galaxy instances via the command line.