# An Introduction to GC3Pie Session-based scripts

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#### What is GC3Pie?

#### GC3Pie is ...

- 1. An *opinionated* Python framework for defining and running computational workflows;
- A rapid development toolkit for running user applications on clusters and IaaS cloud resources;
- 3. The worst name ever given to a middleware piece...

As users, you're mostly interested in this part.

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#### Who am I?

Systems administrator and programmer.

At UZH since 2010, first at GC3 then at S3IT.

Developer of GC3Pie since 2010.

# and what about you?

Name ■ Affiliation ■ Interest in GC3Pie? ■ Programming languages? ■ Experience with command-line or terminal commands?

# About this training, 1

- 1. Concepts and glossary
- 2. Usage of a session-based script
- 3. Command-line tools mainly useful for debugging

# About this training, 2

We'd like the training to be as interactive and informal as possible.

If you have a question, just ask - don't wait.

# About this training, 3

#### These slides are available for download from:

https://github.com/uzh/gc3pie/blob/master/docs/ users/tutorial/slides.pdf

# Concepts and glossary

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However, you need to understand the basic concepts ;-)

#### A typical GC3Pie script?

> ./warholize.py uzh-logo.png --watch 1

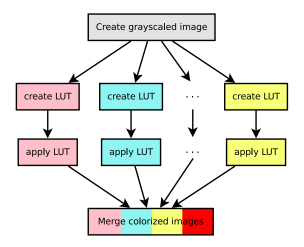






#### How does "Warholize" work?

It is a GC3Pie *workflow*: each box is a distinct application instance that is run.



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# What GC3Pie handles for you

- 1. Resource allocation (e.g. starting new instances on ScienceCloud)
- 2. Selection of resources for each application in the session
- 3. Data transfer (e.g. copying input files in the new instances)
- 4. Remote execution of the application
- 5. Retrieval of results (e.g. copying output files from the running instance)
- 6. De-allocation of resources

# GC3Pie glossary: Application

# GC3Pie runs user applications on clusters and IaaS cloud resources

An Application is just a command to execute.

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An Application is just a command to execute.

If you can run it in the terminal, you can run it in GC3Pie.

# GC3Pie glossary: Application

GC3Pie runs user applications on clusters and IaaS cloud resources

An Application is just a command to execute.

A single execution of an Application is indeed called a Run.

(Other systems might call this a "Job".)

# GC3Pie glossary: Task

GC3Pie runs user applications on clusters and IaaS cloud resources

More generally, GC3Pie runs Tasks.

Tasks are a superset of applications, in that they include workflows.

➤ More on this later!

# GC3Pie glossary: Resources

GC3Pie runs user applications on clusters and IaaS cloud resources

Resources are the computing infrastructures where GC3Pie executes applications.

Resources include: your laptop, the "Hydra" cluster, the Science Cloud, Amazon AWS.

#### Hands-on time!

Start a VM on Science Cloud, using the "GC3Pie Tools Training" snapshot.

#### **Exercise A:**

- 1. Run the ./warholize.py script to get a new "warholized" version of the UZH logo.
- 2. What command-line option lets you run the whole workflow in one go?
- 3. How can you warholize multiple images at once?

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#### Warholize!

> ./warholize.py uzh-logo.png -C 1









# **Session-based scripts**

warholize.py is a typical session-based script.

A session is just a named collection of jobs.

A *session-based script* creates a session and runs all the jobs in it until completion.

# The output directory

If you don't specify an output directory for your job, a session-based script *usually* collects output in the current working directory.

If an output directory already exists, it will be *renamed* and never overwritten.

If you pass the option  $\neg \circ$  directory to the script, all the output dirs will be saved inside that directory.

#### Create a session

A session-based script creates a session and runs all the jobs in it until completion.

Create session logo:

> ./warholize.py uzh-logo.png -s logo

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#### Run a session until done

A session-based script creates a session and runs all the jobs in it until completion.

Run jobs in session logo, polling for updates every 5 seconds:

> ./warholize.py uzh-logo.png -s logo --watch 5

You can stop a GC3Pie script by pressing *Ctrl+C*. Run it again to resume activity from where it stopped.

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A session-based script creates a session and runs all the jobs in it until completion.

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# Alternate display of session contents, I

Display top-level tasks in session logo:

> gsession list logo

**Exercise B:** Now try this yourself.

#### WTF??

In order to work, all GC3Pie utilities need to access the Python script that generated the tasks and session.

To fix: set the PYTHONPATH variable to the directory containing your script:

> export PYTHONPATH=\$PWI

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#### WTF??

```
> gsession list logo
gc3.gc3libs: WARNING: Failed loading file '/home/ubuntu/logo/jobs/WarholizeWorkflow.108': In ...
LoadError: Failed retrieving object from file '/home/ubuntu/logo/jobs/WarholizeWorkflow.108
gc3.gc3libs: WARNING: Ignoring error from loading 'ParallelTaskCollection.107': Failed retr
+-----+
| JobID | Job name | State | Info |
+-----+
+-----+
+-----+
```

In order to work, all GC3Pie utilities need to access the Python script that generated the tasks and session.

To fix: set the PYTHONPATH variable to the directory containing your script:

> export PYTHONPATH=\$PWD

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# Alternate display of session contents, II

#### Display all tasks in session logo:

> gsession list --recursive logo

▶ Workflows and task hierarchy

## Alternate display of session contents, III

Display summary of tasks in session logo:

> gstat -b -s logo

# Display session history

Show log of activity on tasks in session logo:

> gsession log logo

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# **Resource definition**

# The gservers command

The gservers command is used to see configured and available resources.

Resources are defined in file \$HOME/.gc3/gc3pie.conf

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## Example execution resources: local host

Allow GC3Pie to run tasks on the local computer.

This is the default installed by GC3Pie into

\$HOME/.gc3/gc3pie.conf

[resource/localhost]
enabled = yes
type = shellcmd
frontend = localhost
transport = local
max\_cores per\_job = 2
max\_memory\_per\_core = 2GiB
max\_walltime = 12 hours
max\_cores = 2
architecture = x86\_64
auth = none
override = no

## **Example execution resources: SLURM**

Allow submission of jobs to the "Hydra" cluster.

```
[resource/hydra]
enabled = ves
type = slurm
frontend = login.s3it.uzh.ch
transport = ssh
auth = ssh user rmurri
max walltime = 1 day
max cores = 96
max_cores_per_job = 64
max memory_per_core = 1 TiB
architecture = x86 64
prologue_content =
  module load cluster/largemem
[auth/ssh user rmurri]
tvpe=ssh
username=rmurri
```

## Example execution resources: OpenStack

```
# default user on Ubuntu VM images
enabled=no
type=openstack+shellcmd
                                                type=ssh
auth=openstack
                                                username=ubuntu
vm pool_max_size = 32
security group name=default
                                                [auth/openstack]
security group rules=
                                                # only need to set the 'type' here;
 tcp:22:22:0.0.0.0/0,
                                                # any other value will be taken from
 icmp:-1:-1:0.0.0.0/0
                                                # the 'OS *' environment variables
network ids=
                                                type = openstack
 c86b320c-9542-4032-a951-c8a068894cc2
# definition of a single execution VM
```

Allow running tasks on the "ScienceCloud" VM infrastructure

[auth/ssh user ubuntu]

# how to connect
vm\_auth=ssh\_user\_ubuntu
keypair\_name=rmurri
public kev=-/.ssh/id dsa.pub

instance type=1cpu-4ram-hpc

max memory per core = 4 GiB

max cores per job = 8

max\_walltime = 90 days
max cores = 32

architecture = x86 64

image id=2b227d15-8f6a-42b0-b744-ede52ebe59f7

[resource/sciencecloud]

#### Hands-on time!

**Exercise C:** Change the configuration file ~/.gc3/gc3pie.conf to enable the sciencecloud resource. Verify with the gservers command that it works.

#### **Cannot start VMs?**

You need to tell GC3Pie what credentials to use for authenticating to OpenStack / ScienceCloud. You can write them in the configuration file, or use the sc-authenticate.sh script to load them in the environment:

#### > source sc-authenticate.sh

Please enter your ScienceCloud project name [training]:
Please enter your ScienceCloud username [ubuntu]: shortname
Please enter your OpenStack Password:

**Important:** at the "username" prompt, enter your "UZH shortname" – do *not* accept the default!

#### **Select execution resource**

Select where applications will be run with option -r:

```
> ./warholize.py -s logo -r localhost
```

The resource name must exists in the configuration file (i.e., check gservers' output).

Stopping a script and re-starting it with a different resource will likely result in an error: old tasks can no longer be found.

## Passing requirements to the application

Some options are used to specify some requirements of *all* applications in a session:

- -c NUM Set the number of CPU cores required for each job.
  - -m GB Set the amount of memory required per execution core
- -w DURATION Set the time limit for each job; default is script-dependent.

These options have proven not to be much useful except for debugging/experimentation, so they might be removed in a future release!

## Cloud backend management

#### **Hands-on time!**

**Exercise D:** Run the "Warholize" workflow on the ScienceCloud.

Does it succeed? Why?

#### See what VMs are in use

Use the gcloud command to show cloud usage:

> gcloud list

## Limit concurrently-running jobs

Limit the maximum number of concurrently-running tasks with option -J:

```
> ./warholize.py -s logo -J 1
```

In large computational campaigns, it is important not to flood the execution resources with too many tasks.

#### Hands-on time!

#### **Exercise E:**

- 1. Run the "Warholize" workflow on the ScienceCloud. Stop the script while it's running. Now start it again. What happens?
- 2. Run the "Warholize" workflow on the ScienceCloud. Stop the script while it's running; list the jobs in the session and note down the IDs. Now run the script again, adding the "-N" option. When the script terminates, inspect the session again and note the IDs. What has happened? Why?

#### Use -N with caution!

#### From the session-based script's --help output:

-N, --new-session
Discard any information saved in the session dir
(see '--session' option) and start a new session
afresh. Any information about previous jobs is lost.

## Cleanup unused VMs

## Use the gcloud command again:

- To stop *all* unused VMs:
  - > gcloud cleanup

**Exercise F:** Do it. Now.

- To stop a specific VM:
  - > gcloud terminate f031d6ad-bd6c-439e-9a98-4d64

## Session management

## Aborting a single task

To stop and abort a single task, use the gkill command:

> gkill -s logo MyApplication.123

**Exercise:** What happens if you try to abort a task collection?

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To stop and abort a single task, use the gkill command:

> gkill -s logo MyApplication.123

**Exercise:** What happens if you try to abort a task collection?

## Aborting a whole session

Kill all the running tasks in a session again using the gkill command:

> gkill -s logo -A

Alternatively you can use gsession abort:

> gsession abort logo

## Selecting tasks from a session, I

The gselect command is the go-to tool for selective listing of tasks in a session. For example, to list finished tasks:

> gselect -s logo --state TERMINATED

The output of gselect is a list of task IDs, to be fed into another GC3Pie command. For example, to kill all queued tasks:

> gselect -s logo --state SUBMITTED | xargs gkill -s logo

## Selecting tasks from a session, II

The gselect command has many different options to select tasks:

**Exercise:** Use gselect to print the IDs of the "TricolorizeImage" tasks in the last "Warholize" session.

## Thank you!

# Any questions?

GC3Pie manual: http://gc3pie.readthedocs.io/

Mailing-list: gc3pie@googlegroups.com

or read online at

http://dir.gmane.org/gmane.comp.python.gc3pie

## **Workflows**

#### The "Warholize" workflow

How do we "warholize" an arbitrary image?

- 1. Convert the original image to grayscale.
- 2. Colorize the grayscale image using three different colors for each tile.
- 3. Arrange all the colorized images into an  $N \times N$  frame.

**Reference:** http://gc3pie.readthedocs.org/en/master/programmers/tutorials/warholize/warholize.html

## GC3Pie glossary: Task Collections

The basic unit of work in a GC3Pie workflow is called a Task.

The Application class that you already know is a kind of Task (in programming speak, it's a derived class).

A set of Tasks is itself a Task, and is called a TaskCollection.

## Running tasks in sequence

To run tasks in an ordered sequence, one after the other, GC3Pie provides a SequentialTaskCollection class.

It is created with a list of tasks, and runs all of them in the order given. The sequence is dynamic, in that you can add new tasks on the fly, re-run existing ones, or remove future tasks.

A SequentialTaskCollection is itself a task.

## Running tasks in parallel

To run tasks in parallel (i.e., they have no inter-dependency), GC3Pie provides a ParallelTaskCollection class.

It is created with a list of tasks, and runs all of them in parallel (compatibly with the computational resource limits).

A ParallelTaskCollection is itself a task.

## Putting it all together

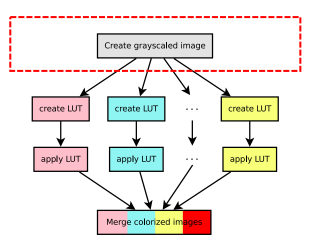
#### So tasks can be:

- Application instances,
- SequentialTaskCollections,
- ParallelTaskCollections.

So you can nest them, and create parallelly-running sequences, or sequences of "job explosions" (many jobs in parallel), or any combination of this.

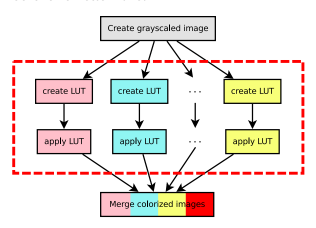
## The Warholize workflow, I

1. Convert the original image to grayscale.



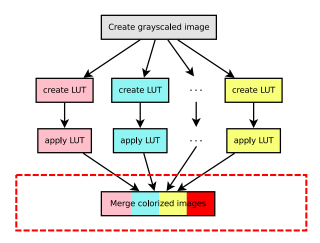
#### The Warholize workflow, II

2. Colorize the grayscale image using three different colors for each tile.



## The Warholize workflow, III

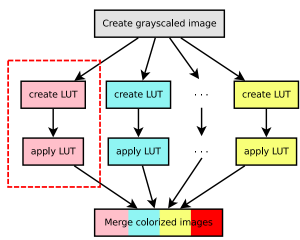
3. Arrange all the colorized images into an  $N \times N$  frame.



## The Warholize workflow, IV

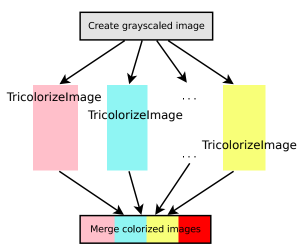
Step 2 actually entails two sub-steps:

- a) mapping greyscale levels to random colors,
- b) applying this mapping to produce new images



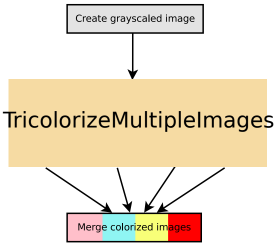
## The Warholize workflow, V

So, Step 2 is a SequentialTaskCollection-type task. Let's call this two-pass sequence TricolorizeImage.



#### The Warholize workflow, VI

All the TricolorizeImage instances run in parallel. Collect them into a ParallelTaskCollection-type task, called TricolorizeMultipleImages.



## The Warholize workflow, VII

Now we are left with a three-step sequence: greyscale, TricolorizeMultipleImages, montage. This can be defined again as a SequentialTaskCollection-type task, the WarholizeWorkflow.



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