# Package 'flowr'

September 25, 2015

| Туре  | Package   |
|-------|---|
| Title | Streamlining Design and Deployment of Complex Workflows   |
| Desci | ription This framework allows you to design and implement complex pipelines, and deploy them on your institution's computing cluster. This has been built keeping in mind the needs of bioinformatics workflows. However, it is easily extendable to any field where a series of steps (shell commands) are to be executed in a (work)flow. |
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| BugR  | Reports https://github.com/sahilseth/flowr/issues   |
| Licen | se MIT + file LICENSE   |
| R to  | opics documented:   |
|       | check check_args fetch flow-class flowopts get_wds job  |

2 check

| Index |                    | 28 |
|-------|--------------------|----|
|       | write_flow_details | 27 |
|       | whisker_render     |    |
|       | verbose            | 26 |
|       | to_flowmat         | 24 |
|       | to_flowdet         | 24 |
|       | to_flowdef         | 21 |
|       | to_flow            | 19 |
|       | test_queue         | 19 |
|       | submit_run         | 18 |
|       | submit_job         | 18 |
|       | submit_flow        | 17 |
|       | status             | 16 |
|       | setup              | 15 |
|       | run                | 14 |
|       | rerun              | 13 |
|       | queue-class        | 11 |
|       | plot_flow          |    |
|       | kill               | g  |

check

Check consistency of flowdef and flowmat

# Description

Check consistency of flowdef and flowmat, using various rules.

## Usage

```
check(x, ...)
## S3 method for class 'flowmat'
check(x, ...)
## S3 method for class 'flowdef'
check(x, verbose = get_opts("verbose"), ...)
```

## Arguments

x a flowdef or flowmat object
... Passed onto either check.flowdef OR check.flowmat functions

verbose A numeric value indicating the amount of messages to produce. Values are integers varying from 0, 1, 2, 3, .... Please refer to the verbose page for more details. [get\_opts("verbose")] [1]

check\_args 3

#### **Details**

## A typical output from flowdef with verbose level: 2

```
checking if required columns are present...
checking if resources columns are present...
checking if dependency column has valid names...
checking if submission column has valid names...
checking for missing rows in def...
checking for extra rows in def...
checking submission and dependency types...
jobname prev.sub_type --> dep_type --> sub_type: relationship
1: aln1_a none --> none --> scatter
2: aln2_a scatter --> none --> scatter
3: sampe_a scatter --> serial --> scatter rel: complex one:one
4: fixrg_a scatter --> serial --> scatter rel: complex one:one
5: merge_a scatter --> gather --> serial rel: many:one
6: markdup_a serial --> serial rel: simple one:one
7: target_a serial --> serial --> serial rel: simple one:one
8: realign_a serial --> burst --> scatter rel: one:many
9: baserecalib_a scatter --> serial --> scatter rel: complex one:one
10: printreads_a scatter --> serial --> scatter rel: complex one:one
```

check\_args

Assert none of the arguemnts of a function are null.

## Description

Checks all the arguments in the parent function and makes sure that none of them are NULL

# Usage

```
check_args(ignore, select)
```

#### **Arguments**

ignore optionally ignore a few variables for checking.
select optionally only check a few variables of the function.

#### **Details**

This function has now been moved to params package.

4 fetch

fetch

Two generic functions to search for pipelines and configuration files.

#### **Description**

These functions help in searching for specific files in the user's space.

fetch\_pipes(): Fetches pipelines in the following places, in this specific order:

```
• user's folder: ~/flowr/pipelines
```

• current wd: ./

**NOTE:** If same pipeline is availabe in multiple places; intitutively, one from the later folder would be selected. As such, giving priority to user's home, and current working directories.

fetch\_conf(): Fetches configuration files in ALL of the following places:

• package: conf folders in flowr and ngsflows packages.

• user's folder: ~/flowr/conf folder.

current wd: ./

**NOTE:** This function would greedily return all matching conf files. One would load all of them in the order returned by this functions. If the same variable is repeated in multiple files, value from the later files would replace those formerly defined. Thus ( as explained above ), giving priority to options defined in user's home and current working directories.

By default flowr loads, flowr.conf and ngsflows.conf. See the details sections, for more explanation on this.

## Usage

```
fetch(x, places, urls, verbose = get_opts("verbose"))
fetch_pipes(x, places, last_only = FALSE,
  urls = get_opts("flowr_pipe_urls"), silent = FALSE,
  verbose = get_opts("verbose"), ask = TRUE)
fetch_conf(x = "flowr.conf", places, ...)
```

#### **Arguments**

| X         | fetch_conf search for files ending with .R and .conf respectively.  |
|-----------|---|
| places    | places (paths) to look for files matching the name. Defaults are already defined in the function.   |
| urls      | urls to look for, works well for pipelines [not implemented yet]  |
| verbose   | A numeric value indicating the amount of messages to produce. Values are integers varying from 0, 1, 2, 3, Please refer to the verbose page for more details. [get_opts("verbose")] [1] |
| last_only | fetch_pipes():. If multiple pipelines match the pattern, return the last one. [TRUE]  |
| silent    | fetch_pipes(): logical, be silent even if no such pipeline is available. [FALSE]  |
| ask       | ask before downloading or copying. [not implemented]  |
|           | [not implemented]   |

flow-class 5

#### **Details**

For example flowr has a variable flow\_run\_path where it puts all the execution logs etc. The default value is picked up from packages's internal flowr.conf file. To redefine this value, one could create a new file called ~/flowr/conf/flowr.conf and add a line:

flow\_run\_path<TAB>my\_awesome\_path, where <TAB> is a tab character, since these are tab seperated files.

Also, at any time you can run, load\_conf('super\_specific\_opts.conf'); to load custom options.

#### See Also

flowopts

## **Examples**

```
## let us find a default conf file
conf = fetch_conf("flowr.conf");conf
## load this
load_opts(conf)

## this returns a list, which prints pretty
pip = fetch_pipes("sleep_pipe")
pip$name
pip$pipe
pip$def
```

flow-class

Describing the flow class

## **Description**

Internal function (used by to\_flow), which aids in creating a flow object.

#### Usage

```
flow(jobs = list(new("job")), name = "newflow", desc = "my_super_flow",
  mode = c("scheduler", "trigger", "R"),
  flow_run_path = get_opts("flow_run_path"), trigger_path = "",
  flow_path = "", version = "0.0", status = "created", execute = "")
```

## Arguments

| jobs          | list: A list of jobs to be included in this flow  |
|---------------|---|
| name          | character: Name of the flow. ['newflow']  |
| desc          | character Description of the flow, used to uniquely identify a flow instance. ['my_super_flow'] |
| mode          | character Mode of submission of the flow (depreciated). ['scheduler']                           |
| flow_run_path | The base path of all the flows you would submit. [~/flows]                                      |
| trigger_path  | character[~/flows/trigger].   |

6 flowopts

```
flow_path character: A unique path identifying a flow instance, populated by submit_flow.

version version of flowr used to create and execute this flow.

status character: Status of the flow.

execute execution status of flow object. [FALSE]
```

#### **Examples**

```
cmds = rep("sleep 5", 10)
qobj <- queue(platform='torque')</pre>
## run the 10 commands in parallel
jobj1 <- job(q_obj=qobj, cmd = cmds, submission_type = "scatter", name = "job1")</pre>
## run the 10 commands sequentially, but WAIT for the previous job to complete
## Many-To-One
jobj2 <- job(q_obj=qobj, cmd = cmds, submission_type = "serial",</pre>
 dependency_type = "gather", previous_job = "job1", name = "job2")
## As soon as first job on 'job1' is complete
## One-To-One
jobj3 <- job(q_obj=qobj, cmd = cmds, submission_type = "scatter",</pre>
 dependency_type = "serial", previous_job = "job1", name = "job3")
fobj <- flow(jobs = list(jobj1, jobj2, jobj3))</pre>
## plot the flow
plot_flow(fobj)
## Not run:
## dry run, only create the structure without submitting jobs
submit_flow(fobj)
## execute the jobs: ONLY works on computing cluster, would fail otherwise
submit_flow(fobj, execute = TRUE)
## End(Not run)
```

flowopts

Default options/params used in flowr and ngsflows

#### **Description**

There are three helper functions which attempt to manage parameters used by flowr and ngsflows:

- get\_opts OR opts\_flow\$get(): show all default options
- set\_opts OR opts\_flow\$set(): set default options
- load\_opts OR opts\_flow\$load(): load options specified in a tab seperated text file

For more details regarding these funtions refer to params package.

```
flowopts

get_opts(...)
set_opts(...)
load_opts(...)
```

flowopts 7

#### **Arguments**

- get: names of options to fetch
- set: a set of options in a name=value format seperated by commas

#### **Format**

```
opts_flow
```

#### **Details**

By default flowr loads, ~/flowr/conf/flowr.conf and ~/flowr/conf/ngsflows.conf Below is a list of default flowr options, retrieved via opts\_flow\$get():

```
Iname
                 |value
|:----|
|default_regex
                 |(.*)
|flow_base_path
                 |~/flowr
|flow_conf_path
               |~/flowr/conf
|flow_parse_lsf |.*(\<[0-9]*\>).*
|flow_parse_moab |(.*)
|flow_parse_sge
                 |(.*)|
|flow_parse_slurm |(.*)
|flow_parse_torque |(.?)\..*
|flow_pipe_paths |~/flowr/pipelines
|flow_pipe_urls
                 |~/flowr/pipelines
|flow_platform
                 llocal
                 |~/flowr/runs
|flow_run_path
|my_conf_path
                 |~/flowr/conf
|my_dir
                 |path/to/a/folder
|my_path
                 |~/flowr
|my_tool_exe
                 |/usr/bin/ls
|time_format
                 |%a %b %e %H:%M:%S CDT %Y |
|verbose
                 | FALSE
```

#### See Also

fetch params read\_sheet

## **Examples**

```
## Set options: set_opts()
opts = set_opts(flow_run_path = "~/mypath")
## OR if you would like to supply a long list of options:
opts = set_opts(.dots = list(flow_run_path = "~/mypath"))
## load options from a configuration file: load_opts()
conffile = fetch_conf("flowr.conf")
load_opts(conffile)
## Fetch options: get_opts()
get_opts("flow_run_path")
get_opts()
```

8 job

| get | wds |
|-----|-----|

Get all the (sub)directories in a folder

#### **Description**

Get all the (sub)directories in a folder

## Usage

```
get_wds(x)
```

#### **Arguments**

Х

path to a folder

job

Describing details of the job object

## **Description**

Internal function (used by to\_flow), which aids in creating a job object.

#### Usage

```
job(cmds = "", name = "myjob", q_obj = new("queue"), previous_job = "",
  cpu = 1, memory, walltime, submission_type = c("scatter", "serial"),
  dependency_type = c("none", "gather", "serial", "burst"), ...)
```

#### **Arguments**

| cmds  | the commands to run |
|-------|---------------------|
| name  | name of the job     |
| q_obj | queue object        |
|       |                     |

previous\_job character vector of previous job. If this is the first job, one can leave this empty,

NA, NULL, '.', or ". In future this could specify multiple previous jobs.

cpu no of cpu's reserved

memory The amount of memory reserved. Units depend on the platform used to process

jobs

walltime The amount of time reserved for this job. Format is unique to a platform. Typi-

cally it looks like 12:00 (12 hours reserved, say in LSF), in Torque etc. we often

see measuring in seconds: 12:00:00

submission\_type

submission type: A character with values: scatter, serial. Scatter means all the 'cmds' would be run in parallel as seperate jobs. Serial, they would combined

into a single job and run one-by-one.

dependency\_type

depedency type. One of none, gather, serial, burst. If previous\_job is specified,

then this would not be 'none'. [Required]

... other passed onto object creation. Example: memory, walltime, cpu

kill 9

#### **Examples**

```
qobj <- queue(platform="torque")</pre>
## torque job with 1 CPU running command 'sleep 2'
jobj <- job(q_obj=qobj, cmd = "sleep 2", cpu=1)</pre>
## multiple commands
cmds = rep("sleep 5", 10)
## run the 10 commands in parallel
jobj1 <- job(q_obj=qobj, cmd = cmds, submission_type = "scatter", name = "job1")</pre>
\#\# run the 10 commands sequentially, but WAIT for the previous job to complete
jobj2 <- job(q_obj=qobj, cmd = cmds, submission_type = "serial",</pre>
   dependency_type = "gather", previous_job = "job1")
fobj <- flow(jobs = list(jobj1, jobj2))</pre>
## plot the flow
plot_flow(fobj)
## Not run:
## dry run, only create the structure without submitting jobs
submit_flow(fobj)
## execute the jobs: ONLY works on computing cluster, would fail otherwise
submit_flow(fobj, execute = TRUE)
## End(Not run)
```

kill

Kill all jobs submitted to the computing platform, for one or multiple flows

## Description

NOTE:

This requires files which are created at the end of the submit\_flow command.

Even if you want to kill the flow, its best to let submit\_flow do its job, when done simply use kill(flow\_wd). If submit\_flow is interrupted, files like flow\_details.rds etc are not created, thus flowr looses the association of jobs with flow instance and cannot monitor, kill or re-run the flow.

```
kill(x, ...)
## S3 method for class 'character'
kill(x, force = FALSE, ...)
## S3 method for class 'flow'
kill(x, kill_cmd, verbose = get_opts("verbose"),
    jobid_col = "job_sub_id", ...)
```

10 plot\_flow

## **Arguments**

| X         | either path to flow wd or object of class flow  |
|-----------|---|
|           | not used  |
| force     | You need to set force=TRUE, to kill multiple flows. This makes sure multiple flows are NOT killed by accident.  |
| kill_cmd  | The command used to kill. flowr tries to guess this commands, as defined in the detect_kill_cmd(). Supplying it here; fot custom platoforms.  |
| verbose   | A numeric value indicating the amount of messages to produce. Values are integers varying from 0, 1, 2, 3, Please refer to the verbose page for more details. [get_opts("verbose")] [1] |
| jobid_col | Advanced use. The column name in 'flow_details.txt' file used to fetch jobids to kill   |

## **Examples**

```
## Not run:
## example for terminal
## flowr kill_flow x=path_to_flow_directory
## In case path matches multiple folders, flowr asks before killing
kill(x='fastq_haplotyper*')
Flowr: streamlining workflows
found multiple wds:
/fastq_haplotyper-MS132-20150825-16-24-04-0Lv1PbpI
/fastq_haplotyper-MS132-20150825-17-47-52-5vFIkrMD
Really kill all of them ? kill again with force=TRUE
## submitting again with force=TRUE will kill them:
kill(x='fastq_haplotyper*', force = TRUE)
## End(Not run)
```

plot\_flow

Plot a clean and scalable flowchart describing the (work)flow

## Description

Plot a flowchart using a flow object or flowdef

```
plot_flow(x, ...)
## S3 method for class 'flow'
plot_flow(x, ...)
## S3 method for class 'list'
plot_flow(x, ...)
## S3 method for class 'character'
```

queue-class 11

```
plot_flow(x, ...)
## S3 method for class 'flowdef'
plot_flow(x, detailed = TRUE, type = c("1", "2"),
   pdf = FALSE, pdffile, ...)
```

## Arguments

| X        | Object of class flow, or a list of flow objects or a flowdef               |
|----------|--|
|          | experimental and only for advanced use.                                    |
| detailed | include submission and dependency types in the plot [TRUE]                 |
| type     | 1 is original, and 2 is a elipse with less details [1]                     |
| pdf      | create a pdf instead of plotting interactively [FALSE]                     |
| pdffile  | <pre>output file name for the pdf file. [flow_path/flow_details.pdf]</pre> |

#### **Examples**

```
qobj = queue(type="lsf")
cmds = rep("sleep 5", 10)
jobj1 <- job(q_obj=qobj, cmd = cmds, submission_type = "scatter", name = "job1")</pre>
jobj2 <- job(q_obj=qobj, name = "job2", cmd = cmds, submission_type = "scatter",</pre>
              dependency_type = "serial", previous_job = "job1")
fobj <- flow(jobs = list(jobj1, jobj2))</pre>
plot_flow(fobj)
### Gather: many to one relationship
jobj1 \leftarrow job(q_obj=qobj, cmd = cmds, submission_type = "scatter", name = "job1")
jobj2 \leftarrow job(q\_obj=qobj, name = "job2", cmd = cmds, submission\_type = "scatter",
             dependency_type = "gather", previous_job = "job1")
fobj <- flow(jobs = list(jobj1, jobj2))</pre>
plot_flow(fobj)
### Burst: one to many relationship
jobj1 <- job(q_obj=qobj, cmd = cmds, submission_type = "serial", name = "job1")</pre>
jobj2 <- job(q_obj=qobj, name = "job2", cmd = cmds, submission_type = "scatter",</pre>
              dependency_type = "burst", previous_job = "job1")
fobj <- flow(jobs = list(jobj1, jobj2))</pre>
plot_flow(fobj)
```

queue-class

A queue object defines details regarding how a job is submitted

## **Description**

Internal function (used by to\_flow), to define the format used to submit a job.

```
queue(object, platform = c("local", "lsf", "torque", "sge", "moab"),
  format = "", queue = "long", walltime, memory, cpu = 1,
  extra_opts = "", submit_exe, nodes = "1", jobname = "name",
  email = Sys.getenv("USER"), dependency = list(), server = "localhost",
  verbose = FALSE, cwd = "", stderr = "", stdout = "", ...)
```

12 queue-class

#### **Arguments**

object this is not used currenlty, ignore.

platform Required and important. Currently supported values are 'lsf' and 'torque'. [Used

by class job]

format [advanced use] We have a default format for the final command line string gen-

erated for 'lsf' and 'torque'.

queue the type of queue your group usually uses 'bsub' etc.

walltime max walltime of a job.

memory The amount of memory reserved. Units depend on the platform used to process

jobs

cpu number of cpus you would like to reserve [Used by class job]

extra\_opts [advanced use] Extra options to be supplied while create the job submission

string.

submit\_exe [advanced use] Already defined by 'platform'. The exact command used to

submit jobs to the cluster example 'qsub'

nodes [advanced use] number of nodes you would like to request. Or in case of torque

name of the nodes. optional [Used by class job]

jobname [debug use] name of this job in the computing cluster

email [advanced use] Defaults to system user, you may put you own email though may

get tons of them.

dependency [debug use] a list of jobs to complete before starting this one

server [not used] This is not implemented currently. This would specify the head node

of the computing cluster. At this time submission needs to be done on the head

node of the cluster where flow is to be submitted

verbose [logical] TRUE/FALSE

cwd [debug use] Ignore stderr [debug use] Ignore stdout [debug use] Ignore

... other passed onto object creation. Example: memory, walltime, cpu

#### **Details**

**Resources**: Can be defined \*\*once\*\* using a queue object and recylced to all the jobs in a flow. If resources (like memory, cpu, walltime, queue) are supplied at the job level they overwrite the one supplied in queue Nodes: can be supplied of extend a job across multiple nodes. This is purely experimental and not supported.

**Server**: This a hook which may be implemented in future.

**Submission script** The 'platform' variable defines the format, and submit\_exe; however these two are avaible for someone to create a custom submission command.

#### **Examples**

```
qobj <- queue(platform='lsf')</pre>
```

rerun 13

rerun Re-run a pipeline in case of hardware or software failures.

## **Description**

- hardware no change required, simple rerun: rerun(x=flow\_wd)
- software either a change to flowmat or flowdef has been made: rerun(x=flow\_wd, mat = new\_flowmat, def = r

#### NOTE:

flow\_wd: flow working directory, same input as used for status

#### Usage

```
rerun(x, ...)
## S3 method for class 'character'
rerun(x, ...)
## S3 method for class 'flow'
rerun(x, mat, def, start_from, execute = TRUE, kill = TRUE,
    select, ignore, verbose = get_opts("verbose"), ...)
```

# Arguments

| x          | flow working directory  |
|------------|---|
|            | passed onto to_flow   |
| mat        | (optional) flowmat fetched from previous submission if missing. For more information regarding the format refer to to_flowmat   |
| def        | (optional) flowdef fetched from previous submission if missing. For more information regarding the format refer to to_flowdef   |
| start_from | which job to start from, this is a job name.  |
| execute    | [logical] whether to execute or not   |
| kill       | (optional) logical indicating whether to kill the jobs from the previous execution of flow.   |
| select     | select a subset of jobs to rerun [character vector]   |
| ignore     | ignore a subset of jobs to rerun [character vector]   |
| verbose    | A numeric value indicating the amount of messages to produce. Values are integers varying from 0, 1, 2, 3, Please refer to the verbose page for more details. [get_opts("verbose")] [1] |

## **Details**

This function fetches details regarding the previous execution from the flow working directory (flow\_wd).

It reads the flow object from the flow\_details.rds file, and extracts flowdef and flowmat from it using to\_flowmat and to\_flowdef functions.

Using new flowmat OR flowdef for re-run:

14 run

Optionally, if either of flowmat or flowdef are supplied; supplied ones are used instead of those extracted from previous submission.

This functions efficiently updates job details of the latest submission into the previous file; thus information regarding previous job ids and their status is not lost.

## **Examples**

```
## Not run:
rerun_flow(wd = wd, fobj = fobj, execute = TRUE, kill = TRUE)
## End(Not run)
```

run

Run automated Pipelines

## Description

Run complete pipelines, by wrapping several steps into one convinient function:

Taking sleep\_pipe as a example.

- Use fetch\_pipes to get paths to a Rscript, flowdef file and optionally a configuration file with various default options used.
- Create a flowmat (using the function defined in the Rscript)
- Create a 'flow' object, using flowmat created and flowdef (as fetched using fetch\_pipes)
- Submit the flow to the cluster (using submit\_flow)

## Usage

```
run(x, platform, def, flow_run_path = get_opts("flow_run_path"),
    execute = FALSE, ...)

run_pipe(x, platform, def, flow_run_path = get_opts("flow_run_path"),
    execute = FALSE, ...)
```

## Arguments

name of the pipeline to run. This is a function called to create a flow\_mat.

platform what platform to use, overrides flowdef

def flow definition

flow\_run\_path passed onto to\_flow. Default it picked up from flowr.conf. Typically this is ~/flowr/runs

execute TRUE/FALSE

passed onto the pipeline function as specified in x

setup 15

#### **Examples**

```
## Not run:
## Run a short pipeline (dry run)
run("sleep_pipe")
## Run a short pipeline on the local machine
run("sleep_pipe", platform = "local", execute = TRUE)
## Run a short pipeline on the a torque cluster (qsub)
run("sleep_pipe", platform = "torque", execute = TRUE)
## Run a short pipeline on the a MOAB cluster (msub)
run("sleep_pipe", platform = "moab", execute = TRUE)
## Run a short pipeline on the a IBM (LSF) cluster (bsub)
run("sleep_pipe", platform = "lsf", execute = TRUE)
## Run a short pipeline on the a MOAB cluster (msub)
run("sleep_pipe", platform = "moab", execute = TRUE)
## change parameters of the pipeline
\ensuremath{\mbox{\#\#}} All extra parameters are passed on to the function function.
run("sleep\_pipe", platform = "lsf", execute = TRUE, x = 5)
## End(Not run)
```

setup

Setup and initialize flowr

## **Description**

This functions creates a directory structure in user's home directory. Additionally it creates a short-cut to the flowr helper script in ~/bin.

## Usage

```
setup(bin = "~/bin", flow_base_path = get_opts("flow_base_path"),
  flow_run_path = get_opts("flow_run_path"),
  flow_conf_path = get_opts("flow_conf_path"),
  flow_pipe_path = get_opts("flow_pipe_paths"))
```

## **Arguments**

```
path to bin folder

flow_base_path The base of flowr configuration and execution folders.

flow_run_path Path to a folder. Main operating folder for this flow. [get_opts("flow_run_path")]
[~/flowr/runs].

flow_conf_path Flowr configuration folder, used by fetch_conf.

flow_pipe_path Folder with all pipelines, used by fetch_pipes.
```

16 status

#### **Details**

Will add more to this, to identify cluster and aid in other things.

status Monitor status of flow(s)

## **Description**

Summarize status of a flow OR multiple flows OR a high-level summary of all flows in a folder.

## Usage

```
status(x, use_cache = FALSE, verbose = get_opts("verbose"),
   out_format = "markdown")

get_status(x, ...)

## S3 method for class 'flow'
get_status(x, verbose, use_cache, out_format, ...)

## S3 method for class 'character'
get_status(x, verbose, use_cache, out_format, ...)

## S3 method for class 'data.frame'
get_status(x, verbose, use_cache, ...)
```

#### **Arguments**

| X          | path to the flow root folder or a parent folder to summarize several flows.   |
|------------|---|
| use_cache  | This skips checking status of jobs which have already been completed a and assumes no new jobs were submitted in the flow(s) being monitored. [FALSE]                                   |
| verbose    | A numeric value indicating the amount of messages to produce. Values are integers varying from 0, 1, 2, 3, Please refer to the verbose page for more details. [get_opts("verbose")] [1] |
| out_format | passed onto knitr:::kable. supports: markdown, rst, html [markdown]   |
|            | not used  |

## **Details**

basename(x) is used in a wild card search.

- Get status of all the flows: (all flows with 'sleep\_pipe' in their name are checked and their status is shown) <br/> flowr status x=~/flowr/runs/sleep\_pipe\*
- Provide a high level summary of ALL flows in a folder: <br/>br> flowr status x=~/flowr/runs

Use **use\_cache=**TRUE to speed up checking the status. This assumes that no new jobs have been submitted and skips (re-)checking status of completed jobs.

Once all the jobs have been submitted to the cluster you may always use use\_cache=TRUE.

submit\_flow 17

#### **Examples**

```
## Not run:
status(x = "~/flowr/runs/sleep_pipe*")
## an example for running from terminal
flowr status x=path_to_flow_directory
## End(Not run)
```

submit\_flow

Submit a flow to the cluster

## **Description**

Submit a flow to the cluster or perform a dry-run to check and debug issues.

#### Usage

```
submit_flow(x, verbose = get_opts("verbose"), ...)
## S3 method for class 'list'
submit_flow(x, verbose = get_opts("verbose"), ...)
## S3 method for class 'flow'
submit_flow(x, verbose = get_opts("verbose"),
    execute = FALSE, uuid, plot = TRUE, dump = TRUE, .start_jid = 1, ...)
```

## **Arguments**

| X          | a object of class flow.   |
|------------|---|
| verbose    | logical.  |
|            | Advanced use. Any additional parameters are passed on to $submit\_job$ function.  |
| execute    | logical whether or not to submit the jobs   |
| uuid       | character Advanced use. This is the final path used for flow execution. Especially useful in case of re-running a flow. |
| plot       | logical whether to make a pdf flow plot (saves it in the flow working directory).                                       |
| dump       | dump all the flow details to the flow path  |
| .start_jid | Job to start this submission from. Advanced use, should be 1 by default.  |

## **Details**

NOTE: Even if you want to kill the flow, its best to let submit\_flow do its job, when done simply use kill(flow\_wd). If submit\_flow is interrupted, files like flow\_details.rds etc are not created, thus flowr looses the association of jobs with flow instance and cannot monitor, kill or re-run the flow.

## **Examples**

```
## Not run:
submit_flow(fobj = fobj, ... = ...)
## End(Not run)
```

18 submit\_run

 $submit_job$ 

Submit a step of a flow

#### **Description**

Internal function (used by submit\_flow), which submit a single step of a flow.

## Usage

```
submit_job(jobj, fobj, job_id, execute = FALSE, verbose = FALSE, ...)
```

## **Arguments**

```
jobj Object of calls job

fobj Object of calls flow

job_id job id

execute A logical vector suggesting whether to submit this job

verbose logical

... not used
```

## **Examples**

```
## Not run:
submit_job(jobj = jobj, fobj = fobj, execute = FALSE,
verbose = TRUE, wd = wd, job_id = job_id)
## End(Not run)
```

submit\_run

Submit several flow objects, limit the max running concurrently

## **Description**

Submit several flow objects, limit the max running concurrently

# Usage

```
submit_run(x, wd, max_processing = 7)
```

# **Arguments**

```
x a list of flow objectswd a folder to monitor (flow_run_path)max_processing max number of flow which may be processed concurrently
```

test\_queue 19

# Description

This function attempts to test the submission of a job to the queue. We would first submit one single job, then submit another with a dependency to see if configuration works. This would create a folder in home called 'flows'.

[Depreciated]: This function has been superseded by run("sleep\_pipe", platform = "lsf", execute=TRUE)

#### Usage

```
test_queue(q_obj, verbose = TRUE, ...)
```

## **Arguments**

```
q_objqueue objectverbosetoggleThese params are passed onto queue. ?queue, for more information
```

## **Examples**

```
## Not run:
test_queue(q_obj = q_obj, ... = ...)
## End(Not run)
```

to\_flow

Create flow objects

#### **Description**

Use a set of shell commands and flow definiton to create flow object.

vector: a file with flowmat table

a named list of commands for a sample. Its best to supply a flowmat instead.

```
to_flow(x, ...)
is.flow(x)

## S3 method for class 'character'
to_flow(x, def, grp_col, jobname_col, cmd_col, ...)

## S3 method for class 'flowmat'
to_flow(x, def, grp_col, jobname_col, cmd_col, flowname,
    flow_run_path, platform, submit = FALSE, execute = FALSE, qobj,
    verbose = get_opts("verbose"), ...)
```

20 to\_flow

```
## $3 method for class 'data.frame'
to_flow(x, ...)
## $3 method for class 'list'
to_flow(x, def, flowname, flow_run_path, desc, qobj,
   verbose = get_opts("verbose"), ...)
```

## **Arguments**

| X             | path (char. vector) to flow_mat, a data.frame or a list.  |  |
|---------------|---|--|
|               | Supplied to specific functions like to_flow.data.frame  |  |
| def           | A flow definition table. Basically a table with resource requirements and mapping of the jobs in this flow  |  |
| grp_col       | column name used to split x (flow_mat). [samplename]  |  |
| jobname_col   | column name with job names. [jobname]   |  |
| cmd_col       | column name with commands. [cmd]  |  |
| flowname      | name of the flow [flowname]   |  |
| flow_run_path | Path to a folder. Main operating folder for this flow. [get_opts("flow_run_path")] [~/flowr/runs].  |  |
| platform      | character vector, specifying the platform to use. local, lsf, torque, moab, sge, slurm, This over-rides the platform column in flowdef. (optional)                                      |  |
| submit        | Use submit_flow on flow object this function returns. TRUE/FALSE. [FALSE]   |  |
| execute       | Use submit_flow on flow object this function returns. TRUE/FALSE, an paramter to submit_flow(). [FALSE]   |  |
| qobj          | Depreciated, modify cluster templates instead. A object of class queue.   |  |
| verbose       | A numeric value indicating the amount of messages to produce. Values are integers varying from 0, 1, 2, 3, Please refer to the verbose page for more details. [get_opts("verbose")] [1] |  |
| desc          | Advanced Use. final flow name, please don't change.   |  |

## **Details**

The parameter x can be a path to a flow\_mat, or a data.frame (as read by read\_sheet). This is a minimum three column matrix with three columns: samplename, jobname and cmd

#### Value

Returns a flow object. If execute=TRUE, fobj is rich with information about where and how the flow was executed. It would include details like jobids, path to exact scripts run etc. To use kill\_flow, to kill all the jobs one would need a rich flow object, with job ids present.

Behaviour: What goes in, and what to expect in return?

- submit=FALSE & execute=FALSE: Create and return a flow object
- submit=TRUE & execute=FALSE: dry-run, Create a flow object then, create a structured execution folder with all the commands
- submit=TRUE, execute=TRUE: Do all of the above and then, submit to cluster

to\_flowdef 21

#### See Also

to\_flowmat, to\_flowdef, to\_flowdet, flowopts and submit\_flow

#### **Examples**

```
## Use this link for a few elaborate examples:
## http://docs.flowr.space/flowr/tutorial.html#define_modules
ex = file.path(system.file(package = "flowr"), "pipelines")
flowmat = as.flowmat(file.path(ex, "sleep_pipe.tsv"))
flowdef = as.flowdef(file.path(ex, "sleep_pipe.def"))
fobj = to_flow(x = flowmat, def = flowdef, flowname = "sleep_pipe", platform = "lsf")
## create a vector of shell commands
cmds = c("sleep 1", "sleep 2")
## create a named list
lst = list("sleep" = cmds)
## create a flowmat
flowmat = to_flowmat(lst, samplename = "samp")
## Use flowmat to create a skeleton flowdef
flowdef = to_flowdef(flowmat)
## use both (flowmat and flowdef) to create a flow
fobj = to_flow(flowmat, flowdef)
## submit the flow to the cluster (execute=TRUE) or do a dry-run (execute=FALSE)
## Not run:
fobj2 = submit_flow(fobj, execute=FALSE)
fobj3 = submit_flow(fobj, execute=TRUE)
## Get the status or kill all the jobs
status(fobj3)
kill(fobj3)
## End(Not run)
```

to\_flowdef

Flow Definition defines how to stich pieces of the (work)flow into a flow.

#### **Description**

This function enables creation of a skeleton flow definition with several default values, using a flowmat. To customize the flowdef, one may supply parameters such as sub\_type and dep\_type upfront. As such, these params must be of the same length as number of unique jobs using in the flowmat.

Each row in this table refers to one step of the pipeline. It describes the resources used by the step and also its relationship with other steps, especially, the step immediately prior to it. <br/> <br/> tr>

**Submission types:** This refers to the sub\_type column in flow definition.<br

to\_flowdef

Consider an example with three steps A, B and C. A has 10 commands from A1 to A10, similarly B has 10 commands B1 through B10 and C has a single command, C1. Consider another step D (with D1-D3), which comes after C.

```
step (number of sub-processes) A (10) \longrightarrow B (10) \longrightarrow C (1) \longrightarrow D (3)
```

- scatter: submit all commands as parallel, independent jobs. Submit A1 through A10 as independent jobs
- serial: run these commands sequentially one after the other.
  - Wrap A1 through A10, into a single job.

#### **Dependency types**

This refers to the dep\_type column in flow definition.

- none: independent job.
  - Initial step A has no dependency
- serial: one to one relationship with previous job.
  - B1 can start as soon as A1 completes, and B2 starts just after A2 and so on.
- gather: *many to one*, wait for **all** commands in the previous job to finish then start the current step.
  - All jobs of B (1-10), need to complete before C1 starts
- burst: *one to many* wait for the previous step which has one job and start processing all cmds in the current step.
  - D1 to D3 are started as soon as C1 finishes.

#### Usage

```
to_flowdef(x, ...)
## S3 method for class 'flowmat'
to_flowdef(x, sub_type, dep_type, prev_jobs,
    queue = "short", platform = "torque", memory_reserved = "2000",
    cpu_reserved = "1", walltime = "1:00", verbose = get_opts("verbose"),
    ...)
## S3 method for class 'flow'
to_flowdef(x, ...)
## S3 method for class 'character'
to_flowdef(x, ...)
as.flowdef(x, ...)
```

#### **Arguments**

```
    can a path to a flowmat, flowmat or flow object.
    not used
    sub_type
    submission type, one of: scatter, serial. Character, of length one or same as the number of jobnames
```

to\_flowdef 23

dep\_type dependency type, one of: gather, serial or burst. Character, of length one or

same as the number of jobnames

prev\_jobs previous job name

queue Cluster queue to be used

platform of the cluster: lsf, sge, moab, torque, slurm etc.

memory\_reserved

amount of memory required.

cpu\_reserved number of cpu's required
walltime amount of walltime required

verbose A numeric value indicating the amount of messages to produce. Values are

integers varying from 0, 1, 2, 3, .... Please refer to the verbose page for more

details. [get\_opts("verbose")] [1]

#### **Format**

This is a tab separated file, with a minimum of 4 columns:<br/>br>

required columns:<br>

• jobname: Name of the step

- sub\_type: Short for submission type, refers to, how should multiple commands of this step be submitted. Possible values are 'serial' or 'scatter'.
- prev\_jobs: Short for previous job, this would be the jobname of the previous job. This can be NA/./none if this is a independent/initial step, and no previous step is required for this to start. Additionally, one may use comma(s) to define multiple previous jobs (A,B).
- dep\_type: Short for dependency type, refers to the relationship of this job with the one defined in 'prev\_jobs'. This can take values 'none', 'gather', 'serial' or 'burst'.

resource columns (recommended):<br>

Additionally, one may customize resource requirements used by each step. The format used varies and depends to the computing platform. Thus its best to refer to your institutions guide to specify these.

- cpu\_reserved integer, specifying number of cores to reserve [1]
- memory\_reserved Usually in KB [2000]
- nodes number of server nodes to reserve, most tools can only use multiple cores on a **single** node [1]
- walltime maximum time allowed for a step, usually in a HH:MM or HH:MM:SS format. [1:00]
- queue the queue to use for job submission [short]

24 to\_flowmat

to\_flowdet

Create a flow's submission detail file

## Description

Create a file describing details regarding jobs ids, submission scripts etc.

## Usage

```
to_flowdet(x, ...)
## S3 method for class 'rootdir'
to_flowdet(x, ...)
## S3 method for class 'character'
to_flowdet(x, ...)
## S3 method for class 'flow'
to_flowdet(x, ...)
```

## **Arguments**

x this is a wd ... not used

#### **Details**

The path provided should contain a flow\_detaits.rds file (which is used to extract all the information).

Incase a parent folder with multiple flows is provided information regarding jobids is omitted.

if x is char. assumed a path, check if flow object exists in it and read it. If there is no flow object, try using a simpler function

to\_flowmat

Create a flowmat using a list a commands.

## Description

Create a flowmat (data.frame) using a named list a commands.

as.flowmat(): reads a file and checks for required columns. If x is data.frame checks for required columns.

to\_flowmat 25

#### Usage

```
to_flowmat(x, ...)
## S3 method for class 'list'
to_flowmat(x, samplename, ...)
## S3 method for class 'data.frame'
to_flowmat(x, ...)
## S3 method for class 'flow'
to_flowmat(x, ...)
as.flowmat(x, grp_col, jobname_col, cmd_col, ...)
is.flowmat(x)
```

#### **Arguments**

a named list, where name corresponds to the jobname and value is a vector of commands to run.
 not used
 character of length 1 or that of nrow(x) ['samplename']
 column used for grouping, default samplename.
 jobname\_col
 column specifying jobname, default jobname
 column specifying commands to run, default cmd

#### **Examples**

```
## Use this link for a few examples:
## http://docs.flowr.space/flowr/tutorial.html#define_modules
## create a vector of shell commands
cmds = c("sleep 1", "sleep 2")
## create a named list
lst = list("sleep" = cmds)
## create a flowmat
flowmat = to_flowmat(lst, samplename = "samp")
## Use flowmat to create a skeleton flowdef
flowdef = to_flowdef(flowmat)
## use both (flowmat and flowdef) to create a flow
fobj = to_flow(flowmat, flowdef)
## submit the flow to the cluster (execute=TRUE) or do a dry-run (execute=FALSE)
## Not run:
fobj2 = submit_flow(fobj, execute=FALSE)
fobj3 = submit_flow(fobj, execute=TRUE)
## Get the status or kill all the jobs
status(fobj3)
kill(fobj3)
## End(Not run)
```

26 verbose

verbose

Verbose levels, defining verboseness of messages

#### **Description**

There are several levels of verboseness one can choose from.

#### levels:

- level 0 is almost silent, producing only necessary messages
- level 1 is good for most purposes, where as,
- level 2 is good when developing a new pipeline.
- level 3 is good for debugging, especially when getting un-expected results.

One can set the level of verboseness using opts\_flow\$set(verbose=2), which will be used across flowr and ngsflows packages. Additionally one may set this value in the configurations files: ~/flowr/conf/flowr.conf OR ~/flowr/conf/ngsflows.conf.

#### Usage

verbose

#### **Format**

NULL

#### **Examples**

```
fl = system.file("pipelines/abcd.def", package = "flowr")
def = as.flowdef(f1, verbose = 0)
# def seems to be a file, reading it...
def = as.flowdef(f1, verbose = 1)
# def seems to be a file, reading it...
# checking if required columns are present...
# checking if resources columns are present...
# checking if dependency column has valid names...
# checking if submission column has valid names...
# checking for missing rows in def...
# checking for extra rows in def...
# checking submission and dependency types...
def = as.flowdef(f1, verbose = 2)
# def seems to be a file, reading it...
# checking if required columns are present...
# checking if resources columns are present...
# checking if dependency column has valid names...
# checking if submission column has valid names...
# checking for missing rows in def...
# checking for extra rows in def...
# checking submission and dependency types...
# jobname prev.sub_type --> dep_type --> sub_type: relationship
  1: A none --> none --> scatter
  2: B scatter --> serial --> scatter rel: complex one:one
  3: C scatter --> gather --> serial rel: many:one
# 4: D serial --> burst --> scatter rel: one:many
```

whisker\_render 27

| whisker | randar |  |
|---------|--------|--|
| wnisker | render |  |

Wrapper around whisker.render with some additional checks

# Description

Internal function (used by submit\_job), which creates a submission script using platform specific templates.

## Usage

```
whisker_render(template, data)
```

## **Arguments**

template template used

data a list with variables to be used to fill in the template.

write\_flow\_details

write files desribing this flow

## Description

write files desribing this flow

## Usage

```
write_flow_details(x, fobj, summ, flow_det, plot = FALSE)
```

# **Arguments**

x path to write to fobj flow object

summ a status summary.

flow\_det a flow details data.frame

plot logical, plot or not

# Index

| *Topic datasets                       | queue, <i>12</i> , <i>20</i>           |
|---------------------------------------|--|
| flowopts, 6                           | queue (queue-class), 11                |
| verbose, 26                           | queue-class, 11                        |
| *Topic <b>queue</b>                   |  |
| queue-class, 11                       | read_sheet, 7                          |
|                                       | rerun, 13                              |
| <pre>as.flowdef(to_flowdef), 21</pre> | run, 14                                |
| as.flowmat(to_flowmat), 24            | run_flow(run),14                       |
|                                       | run_pipe (run), 14                     |
| check, 2                              |  |
| check_args, 3                         | set_opts, 6                            |
|                                       | <pre>set_opts (flowopts), 6</pre>      |
| definition (to_flowdef), 21           | setup, 15                              |
| Catala 4.7                            | status, <i>13</i> , 16                 |
| fetch, 4, 7                           | submit_flow, 6, 9, 14, 17, 21          |
| fetch_conf, 4, 15                     | submit_job, <i>17</i> , 18             |
| fetch_conf (fetch), 4                 | submit_run, 18                         |
| fetch_pipes, 4, 14, 15                |  |
| fetch_pipes (fetch), 4                | test_queue, 19                         |
| flow, 10, 13, 18, 19                  | to_flow, <i>5</i> , <i>11</i> , 19     |
| flow(flow-class), 5                   | to_flowdef, <i>13</i> , <i>21</i> , 21 |
| flow-class, 5                         | to_flowdet, <i>21</i> , 24             |
| flowdef (to_flowdef), 21              | to_flowmat, <i>13</i> , <i>21</i> , 24 |
| flowopts, 5, 6, 21                    |  |
| flowr (to_flow), 19                   | verbose, 2, 4, 10, 13, 16, 20, 23, 26  |
| got onto 6                            | 1                                      |
| get_opts, 6                           | whisker_render, 27                     |
| get_opts (flowopts), 6                | write_flow_details,27                  |
| get_status (status), 16               |  |
| get_wds, 8                            |  |
| is.flow(to_flow), 19                  |  |
| is.flowdef(to_flowdef), 21            |  |
| is.flowmat(to_flowmat), 24            |  |
| 13.110Milat (to_110Milat), 21         |  |
| job, 8, 18                            |  |
|                                       |  |
| kill, 9                               |  |
| 1                                     |  |
| load_opts, 6                          |  |
| <pre>load_opts (flowopts), 6</pre>    |  |
| opts_flow(flowopts), 6                |  |
| ορισ_1 τον (1 τονορισ), θ             |  |
| params, 6, 7                          |  |
| plot_flow, 10                         |  |