batchtools

Contents

Setup	1
Cluster Functions	1
Template Files	2
Configuration File	
Migration from BatchJobs/Batchexperiments	2
Internal Changes	3
Interface Changes	3
Template changes	
New features	
Porting to batchtools	
Example 1: Approximation of π	4
Example 2: Machine Learning	6
Problems and Algorithms	6
Creating jobs	
Before Submitting	
Submitting and Collecting Results	
Example: Error Handling	11
Workflow	13
On the Local System	13
On Multiple Systems	

Setup

Cluster Functions

The communication with the batch system is managed via so-called cluster functions. They are created with the constructor makeClusterFunctions which defines how jobs are submitted on your system. Furthermore, you may provide functions to list queued/running jobs and to kill jobs.

Usually you do not have to start from scratch but can just use one of the cluster functions which ship with the package:

- Interactive Cluster Functions (default): docs, implementation
- Multicore Cluster Functions: docs, implementation
- Socket Cluster Functions: docs, implementation
- Makeshift SSH cluster: docs, implementation
- Docker Swarm: docs, implementation
- IBM Spectrum Load Sharing Facility (LSF): docs, implementation
- OpenLava: docs, implementation
- Univa Grid Engine / Oracle Grid Engine (OGE) / Sun Grid Engine (SGE): docs, implementation
- Slurm: docs, implementation
- TORQUE/OpenPBS: docs, implementation

To use the package with the socket cluster functions, you would call the respective constructor makeCluster-FunctionsSocket():

```
reg = makeRegistry(NA)
reg$cluster.functions = makeClusterFunctionsSocket(2)
```

To make this selection permanent for this registry, save the Registry with saveRegistry(). To make your cluster function selection permanent for a specific system across R sessions for all new Registries, you can set up a configuration file (see below).

If you have trouble debugging your cluster functions, you can enable the debug mode for extra output. To do so, install the debugme package and set the environment variable DEBUGME to batchtools before you load the batchtools package:

```
Sys.setenv(DEBUGME = "batchtools")
library(batchtools)
```

Template Files

Many cluster functions require a template file as argument. These templates are used to communicate with the scheduler and contain placeholders to evaluate arbitrary R expressions. Internally, the brew package is used for this purpose. Some exemplary template files can be found here. It would be great if you would help expand this collection to cover more exotic configurations. To do so, please send your template via mail or open a new pull request.

Note that all variables defined in a JobCollection can be used inside the template. If you need to pass extra variables, you can set them via the argument resources of submitJobs().

If the flexibility which comes with templating is not sufficient, you can still construct a custom cluster function implementation yourself using the provided constructor.

Configuration File

The configuration file can be used to set system specific options. Its default location depends on the operating system (see Registry), but for the first time setup you can put one in the current working directory (as reported by getwd()). In order to set the cluster function implementation, you would generate a file with the following content:

```
cluster.functions = makeClusterFunctionsInteractive()
```

The configuration file is parsed whenever you create or load a Registry. It is sourced inside of your registry which has the advantage that you can (a) access all of the parameters which are passed to makeRegistry and (b) you can also directly change them. Lets say you always want your working directory in your home directory and you always want to load the checkmate package on the nodes, you can just append these lines:

```
work.dir = "~"
packages = union(packages, "checkmate")
```

See the documentation on Registry for a more complete list of supported configuration options.

Migration from BatchJobs/Batchexperiments

The development of BatchJobs and BatchExperiments is discontinued because of the following reasons:

- Maintainability: The packages BatchJobs and BatchExperiments are tightly connected which makes
 maintaining difficult. Changes have to be synchronized and tested against the current CRAN versions
 for compatibility. Furthermore, BatchExperiments violates CRAN policies by calling internal functions
 of BatchJobs.
- Data base issues: Although we invested weeks to mitigate issues with locks of the SQLite data base or file system (staged queries, file system timeouts, ...), BatchJobs kept working unreliable on some systems with high latency or specific file systems. This made BatchJobs unusable for many users.

BatchJobs and BatchExperiments will remain on CRAN, but new features are unlikely to be ported back.

Internal Changes

- batchtools does not use SQLite anymore. Instead, all the information is stored directly in the registry using data.tables acting as an in-memory database. As a side effect, many operations are much faster.
- Nodes do not have to access the registry. submitJobs() stores a temporary object of type JobCollection on the file system which holds all the information necessary to execute a chunk of jobs via doJobCollection() on the node. This avoids file system locks because each job accesses only one file exclusively.
- ClusterFunctionsMulticore now uses the parallel package for multicore execution.
- ClusterFunctionsSSH can still be used to emulate a scheduler-like system which respects the work load on the local machine. Setting the hostname to "localhost" just strips out ssh of the command issued.

Interface Changes

- batchtools remembers the last created or loaded Registry and sets it as default registry. This way, you do not need to pass the registry around anymore. If you need to work with multiple registries simultaneously on the other hand, you can still do so by explicitly passing registries to the functions.
- Most functions now return a data.table which is keyed with the job.id. This way, return values can be joined together easily and efficient (see this help page for some examples).
- The building blocks of a problem has been renamed from static and dynamic to the more intuitive data and fun. Thus, algorithm function should have the formal arguments job, data and instance.
- The function makeDesign has been removed. Parameters can be defined by just passing a data.frame or data.table to addExperiments. For exhaustive designs, use data.table::CJ().

Template changes

• The scheduler should directly execute the command:

```
Rscript -e 'batchtools::doJobCollection(<filename>)'
```

There is no intermediate R source file like there was in BatchJobs.

- $\bullet \ \ {\rm All\ information\ stored\ in\ the\ object\ {\tt JobCollection\ } can\ be\ accessed\ while\ brewing\ the\ template}.$
- Extra variables may be passed via the argument resource of submitJobs.

New features

- Support for Docker Swarm via ClusterFunctionsDocker.
- Jobs can now be tagged and untagged to provide an easy way to group them.
- Some resources like the number of CPUs are now optionally passed to parallelMap. This eases nested parallelization, e.g. to use multicore parallelization on the slave by just setting a resource on the master. See submitJobs() for an example.

- ClusterFunctions are now more flexible in general as they can define hook functions which will be called at certain events. ClusterFunctionsDocker is an example use case which implements a housekeeping routine. This routine is called every time before a job is about to get submitted to the scheduler (in the case: the Docker Swarm) via the hook pre.submit and every time directly after the registry synchronized jobs stored on the file system via the hook post.sync.
- More new features are covered in the NEWS.

Porting to batchtools

The following table assists in porting to batchtools by mapping BatchJobs/BatchExperiments functions to their counterparts in batchtools. The table does not cover functions which are (a) used only internally in BatchJobs and (b) functions which have not been renamed.

BatchJobs	batchtools	
addRegistryPackagesSet reg\$packages or		
	${\tt reg\$namespaces},{\tt call}$	
	saveRegistry()	
${\tt addRegistrySource}$	Dirs -	
$\verb"addRegistrySourceFi" \textbf{Sets} \texttt{ reg} \texttt{\$source}, \ call$		
	saveRegistry()	
${\tt batchExpandGrid}$	batchMap:	
	batchMap(, args	
	= CJ(x = 1:3, y =	
	1:10))	
batchMapQuick	btmapply	
batchReduceResult	s -	
batchUnexport	batchExport	
filterResults	-	
getJobIds	$\operatorname{findJobs}$	
getJobInfo	${\it getJobStatus}$	
getJob	${\it make Job}$	
${\tt getJobParamDf}$	$\operatorname{getJobPars}$	
loadResults	${\bf reduce Results List}$	
$\verb reduceResultsDataFrame ce ResultsDataTable $		
$\verb reduceResultsMatrix reduceResultsList +$		
	<pre>do.call(rbind,</pre>	
	res)	
${\tt reduceResultsVect}$	\mathbf{or} educeResultsDataTable	
${\tt setJobFunction}$	-	
setJobNames	-	
showStatus	$\operatorname{getStatus}$	

Example 1: Approximation of π

To get a first insight into the usage of batchtools, we start with an exemplary Monte Carlo simulation to approximate π . For background information, see Wikipedia.

First, a so-called registry object has to be created, which defines a directory where all relevant information, files and results of the computational jobs will be stored. There are two different types of registry objects: First, a regular Registry which we will use in this example. Second, an ExperimentRegistry which provides an alternative way to define computational jobs and thereby is tailored for a broad range of large scale

computer experiments (see, for example, this vignette). Here, we use a temporary registry which is stored in the temp directory of the system and gets automatically deleted if you close the R session.

```
reg = makeRegistry(file.dir = NA, seed = 1)
```

For a permanent registry, set the file.dir to a valid path. It can then be reused later, e.g., when you login to the system again, by calling the function loadRegistry(file.dir).

When a registry object is created or loaded, it is stored for the active R session as the default. Therefore the argument reg will be ignored in functions calls of this example, assuming the correct registry is set as default. To get the current default registry, getDefaultRegistry can be used. To switch to another registry, use setDefaultRegistry().

First, we create a function which samples n points (x_i, y_i) whereas x_i and y_i are distributed uniformly, i.e. $x_i, y_i \sim \mathcal{U}(0, 1)$. Next, the distance to the origin (0, 0) is calculated and the fraction of points in the unit circle $(d \leq 1)$ is returned.

```
piApprox = function(n) {
   nums = matrix(runif(2 * n), ncol = 2)
   d = sqrt(nums[, 1]^2 + nums[, 2]^2)
   4 * mean(d <= 1)
}
piApprox(1000)</pre>
```

[1] 3.132

We now parallelize piApprox() with batchtools: We create 10 jobs, each doing a MC simulation with 10⁵ jobs. We use batchMap() to define the jobs (note that this does not yet start the calculation):

```
batchMap(fun = piApprox, n = rep(1e5, 10))
```

```
## Adding 10 jobs ...
```

The length of the vector or list defines how many different jobs are created, while the elements itself are used as arguments for the function. The function batchMap(fun, ...) works analogously to Map(f, ...) of the base package. An overview over the jobs and their IDs can be retrieved with getJobTable() which returns a data frame with all relevant information:

names(getJobTable())

```
[1] "job.id"
                         "submitted"
                                         "started"
                                                         "done"
##
    [5] "error"
                         "memory"
                                         "batch.id"
                                                         "log.file"
    [9] "job.hash"
                         "job.name"
                                         "time.queued"
                                                         "time.running"
##
                         "tags"
## [13] "n"
```

Note that a unique job ID is assigned to each job. These IDs can be used to restrict operations to subsets of jobs. To actually start the calculation, call **submitJobs()**. The registry and the selected job IDs can be taken as arguments as well as an arbitrary list of resource requirements, which are to be handled by the cluster back end.

```
submitJobs(resources = list(walltime = 3600, memory = 1024))
```

```
## Submitting 10 jobs in 10 chunks using cluster functions 'Interactive' ...
```

In this example, a cap for the execution time (so-called walltime) and for the maximum memory requirements are set. The progress of the submitted jobs can be checked with getStatus().

```
getStatus()
```

```
## Status for 10 jobs:
## Submitted : 10 (100.0%)
```

```
##
     Started
                : 10 (100.0%)
##
                : 10 (100.0%)
     Done
##
     Error
                   0 (
                        0.0%)
##
                         0.0%)
     Queued
                   0 (
##
     Running
                   0 (
                         0.0%)
                        0.0%)
##
     Expired
                   0 (
```

The resulting output includes the number of jobs in the registry, how many have been submitted, have started to execute on the batch system, are currently running, have successfully completed, and have terminated due to an R exception. After jobs have successfully terminated, we can load their results on the master. This can be done in a simple fashion by using either <code>loadResult()</code>, which returns a single result exactly in the form it was calculated during mapping, or by using <code>reduceResults()</code>, which is a version of <code>Reduce()</code> from the base package for registry objects.

```
waitForJobs()
## [1] TRUE
mean(sapply(1:10, loadResult))
## [1] 3.140652
reduceResults(function(x, y) x + y) / 10
## [1] 3.140652
```

If you are absolutely sure that your function works, you can take a shortcut and use batchtools in an lapply fashion using btlapply(). This function creates a temporary registry (but you may also pass one yourself), calls batchMap(), wait for the jobs to terminate with waitForJobs() and then uses reduceResultsList() to return the results.

```
res = btlapply(rep(1e5, 10), piApprox)
mean(unlist(res))
```

[1] 3.142396

Example 2: Machine Learning

We stick to a rather simple, but not unrealistic example to explain some further functionalities: Applying two classification learners to the famous iris data set (Anderson 1935), vary a few hyperparameters and evaluate the effect on the classification performance.

First, we create a registry, the central meta-data object which records technical details and the setup of the experiments. We use an <code>ExperimentRegistry</code> where the job definition is split into creating problems and algorithms. See the paper on <code>BatchJobs</code> and <code>BatchExperiments</code> for a detailed explanation. Again, we use a temporary registry and make it the default registry.

```
library(batchtools)
reg = makeExperimentRegistry(file.dir = NA, seed = 1)
```

Problems and Algorithms

By adding a problem to the registry, we can define the data on which certain computational jobs shall work. This can be a matrix, data frame or array that always stays the same for all subsequent experiments. But it can also be of a more dynamic nature, e.g., subsamples of a dataset or random numbers drawn from a probability distribution. Therefore the function addProblem() accepts static parts in its data argument,

which is passed to the argument fun which generates a (possibly stochastic) problem instance. For data, any R object can be used. If only data is given, the generated instance is data. The argument fun has to be a function with the arguments data and job (and optionally other arbitrary parameters). The argument job is an object of type Job which holds additional information about the job.

We want to split the iris data set into a training set and test set. In this example we use use subsampling which just randomly takes a fraction of the observations as training set. We define a problem function which returns the indices of the respective training and test set for a split with 100 * ratio% of the observations being in the test set:

```
subsample = function(data, job, ratio, ...) {
  n = nrow(data)
  train = sample(n, floor(n * ratio))
  test = setdiff(seq_len(n), train)
  list(test = test, train = train)
}
```

addProblem() files the problem to the file system and the problem gets recorded in the registry.

```
data("iris", package = "datasets")
addProblem(name = "iris", data = iris, fun = subsample, seed = 42)
```

Adding problem 'iris'

The function call will be evaluated at a later stage on the workers. In this process, the data part will be loaded and passed to the function. Note that we set a problem seed to synchronize the experiments in the sense that the same resampled training and test sets are used for the algorithm comparison in each distinct replication.

The algorithms for the jobs are added to the registry in a similar manner. When using addAlgorithm(), an identifier as well as the algorithm to apply to are required arguments. The algorithm must be given as a function with arguments job, data and instance. Further arbitrary arguments (e.g., hyperparameters or strategy parameters) may be defined analogously as for the function in addProblem. The objects passed to the function via job and data are here the same as above, while via instance the return value of the evaluated problem function is passed. The algorithm can return any R object which will automatically be stored on the file system for later retrieval. Firstly, we create an algorithm which applies a support vector machine:

```
svm.wrapper = function(data, job, instance, ...) {
   library("e1071")
   mod = svm(Species ~ ., data = data[instance$train, ], ...)
   pred = predict(mod, newdata = data[instance$test, ], type = "class")
   table(data$Species[instance$test], pred)
}
addAlgorithm(name = "svm", fun = svm.wrapper)
```

Adding algorithm 'svm'

Secondly, a random forest of classification trees:

```
forest.wrapper = function(data, job, instance, ...) {
   library("ranger")
   mod = ranger(Species ~ ., data = data[instance$train, ], write.forest = TRUE)
   pred = predict(mod, data = data[instance$test, ])
   table(data$Species[instance$test], pred$predictions)
}
addAlgorithm(name = "forest", fun = forest.wrapper)
```

Adding algorithm 'forest'

Both algorithms return a confusion matrix for the predictions on the test set, which will later be used to calculate the misclassification rate.

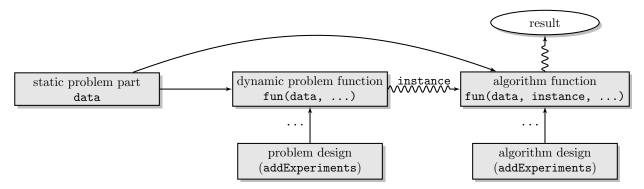
Note that using the ... argument in the wrapper definitions allows us to circumvent naming specific design parameters for now. This is an advantage if we later want to extend the set of algorithm parameters in the experiment. The algorithms get recorded in the registry and the corresponding functions are stored on the file system.

Defined problems and algorithms can be queried:

```
getProblemIds()
## [1] "iris"
getAlgorithmIds()
```

[1] "svm" "forest"

The flow to define experiments is summarized in the following figure:



Creating jobs

addExperiments() is used to parametrize the jobs and thereby define computational jobs. To do so, you have to pass named lists of parameters to addExperiments(). The elements of the respective list (one for problems and one for algorithms) must be named after the problem or algorithm they refer to. The data frames contain parameter constellations for the problem or algorithm function where columns must have the same names as the target arguments. When the problem design and the algorithm design are combined in addExperiments(), each combination of the parameter sets of the two designs defines a distinct job. How often each of these jobs should be computed can be determined with the argument repls.

```
# problem design: try two values for the ratio parameter
pdes = list(iris = data.table(ratio = c(0.67, 0.9)))

# algorithm design: try combinations of kernel and epsilon exhaustively,
# try different number of trees for the forest
ades = list(
    svm = CJ(kernel = c("linear", "polynomial", "radial"), epsilon = c(0.01, 0.1)),
    forest = data.table(ntree = c(100, 500, 1000))
)
addExperiments(pdes, ades, repls = 5)
```

```
## Adding 60 experiments ('iris'[2] x 'svm'[6] x repls[5]) ...
## Adding 30 experiments ('iris'[2] x 'forest'[3] x repls[5]) ...
```

The jobs are now available in the registry with an individual job ID for each. The function summarizeExperiments() returns a table which gives a quick overview over all defined experiments.

```
summarizeExperiments()
##
      problem algorithm .count
##
       <fctr>
                  <fctr>
                          <int>
## 1:
         iris
                             60
                     svm
                             30
## 2:
         iris
                  forest
summarizeExperiments(by = c("problem", "algorithm", "ratio"))
##
      problem algorithm ratio .count
##
       <fctr>
                  <fctr> <num>
                                 <int>
## 1:
                          0.67
         iris
                     svm
                                    30
## 2:
         iris
                     svm 0.90
                                    30
## 3:
         iris
                  forest
                          0.67
                                    15
```

Before Submitting

iris

forest

0.90

15

4:

Before submitting all jobs to the batch system, we encourage you to test each algorithm individually. Or sometimes you want to submit only a subset of experiments because the jobs vastly differ in runtime. Another reoccurring task is the collection of results for only a subset of experiments. For all these use cases, findExperiments() can be employed to conveniently select a particular subset of jobs. It returns the IDs of all experiments that match the given criteria. Your selection can depend on substring matches of problem or algorithm IDs using prob.name or algo.name, respectively. You can also pass R expressions, which will be evaluated in your problem parameter setting (prob.pars) or algorithm parameter setting (algo.pars). The expression is then expected to evaluate to a Boolean value. Furthermore, you can restrict the experiments to specific replication numbers.

To illustrate findExperiments(), we will select two experiments, one with a support vector machine and the other with a random forest and the parameter ntree = 1000. The selected experiment IDs are then passed to testJob.

```
id1 = head(findExperiments(algo.name = "svm"), 1)
print(id1)
##
      job.id
##
       <int>
## 1:
           1
id2 = head(findExperiments(algo.name = "forest", algo.pars = (ntree == 1000)), 1)
print(id2)
##
      job.id
##
       <int>
## 1:
          71
testJob(id = id1)
## Generating problem instance for problem 'iris' ...
## Applying algorithm 'svm' on problem 'iris' ...
##
##
                 setosa versicolor virginica
                     17
                                            0
##
     setosa
                                 0
                                            2
##
     versicolor
                      0
                                 16
```

```
virginica
                                           15
testJob(id = id2)
## Generating problem instance for problem 'iris' ...
## Applying algorithm 'forest' on problem 'iris' ...
##
##
                 setosa versicolor virginica
##
                     17
     setosa
                                  0
##
                      0
                                 16
                                            2
     versicolor
##
     virginica
                      0
                                  1
                                           14
```

If something goes wrong, batchtools comes with a bunch of useful debugging utilities (see separate vignette on error handling). If everything turns out fine, we can proceed with the calculation.

Submitting and Collecting Results

To submit the jobs, we call submitJobs() and wait for all jobs to terminate using waitForJobs().

```
submitJobs()
```

```
## Submitting 90 jobs in 90 chunks using cluster functions 'Interactive' ...
waitForJobs()
```

```
## [1] TRUE
```

After jobs are finished, the results can be collected with reduceResultsDataTable() where we directly extract the mean misclassification error:

```
reduce = function(res) list(mce = (sum(res) - sum(diag(res))) / sum(res))
results = reduceResultsDataTable(fun = reduce)
head(results)
```

```
##
      job.id
               mce
##
       <int> <num>
## 1:
           1 0.04
## 2:
           2 0.00
## 3:
           3 0.06
## 4:
           4 0.04
## 5:
           5 0.02
## 6:
           6
              0.04
```

Next, we merge the results table with the table of job parameters using one of the join helpers provided by batchtools (here, we use an inner join):

```
tab = ijoin(getJobPars(), results)
head(tab)
```

```
##
      job.id problem algorithm ratio kernel epsilon ntree
##
       <int>
              <fctr>
                         <fctr> <num> <char>
                                               <num> <num> <num>
## 1:
           1
                iris
                            svm 0.67 linear
                                                0.01
                                                         NA
                                                             0.04
                                                0.01
## 2:
           2
                iris
                                 0.67 linear
                                                         NA
                                                             0.00
                            svm
## 3:
           3
                                 0.67 linear
                                                0.01
                                                             0.06
                iris
                            svm
                                                         NA
           4
                                                0.01
                                                            0.04
## 4:
                            svm 0.67 linear
                                                         NA
                iris
                            svm 0.67 linear
                                                0.01
                                                         NA 0.02
## 5:
           5
                iris
## 6:
           6
                iris
                            svm 0.67 linear
                                                0.10
                                                         NA 0.04
```

We now aggregate the results group-wise. You can use data.table, base::aggregate(), or the dplyr package for this purpose. Here, we use data.table to subset the table to jobs where the ratio is 0.67 and group by algorithm the algorithm hyperparameters:

```
tab[ratio == 0.67, list(mmce = mean(mce)),
  by = c("algorithm", "kernel", "epsilon", "ntree")]
```

```
##
                     kernel epsilon ntree mmce
      algorithm
##
         <fctr>
                     <char>
                               <num> <num> <num>
## 1:
                                         NA 0.032
                                0.01
             svm
                     linear
## 2:
             svm
                     linear
                                0.10
                                         NA 0.032
## 3:
             svm polynomial
                                0.01
                                         NA 0.088
## 4:
             svm polynomial
                                0.10
                                         NA 0.088
## 5:
                                0.01
                                         NA 0.048
                     radial
             svm
## 6:
                     radial
                                0.10
                                         NA 0.048
             svm
## 7:
                                  NA
                                        100 0.048
         forest
                          NA
## 8:
         forest
                          NA
                                  NA
                                        500 0.052
## 9:
         forest
                          NA
                                  NA
                                       1000 0.044
```

Example: Error Handling

In any large scale experiment many things can and will go wrong. The cluster might have an outage, jobs may run into resource limits or crash, subtle bugs in your code could be triggered or any other error condition might arise. In these situations it is important to quickly determine what went wrong and to recompute only the minimal number of required jobs.

Therefore, before you submit anything you should use testJob() to catch errors that are easy to spot because they are raised in many or all jobs. If external is set, this function runs the job without side effects in an independent R process on your local machine via Rscript similar as on the slave, redirects the output of the process to your R console, loads the job result and returns it. If you do not set external, the job is executed is in the currently running R session, with the drawback that you might be unable to catch missing variable declarations or missing package dependencies.

By way of illustration here is a small example. First, we create a temporary registry.

```
library(batchtools)
reg = makeRegistry(file.dir = NA, seed = 1)
```

Ten jobs are created, one will trow a warning and two of them will raise an exception.

```
flakeyFunction <- function(value) {
  if (value == 5) warning("Just a simple warning")
  if (value %in% c(2, 9)) stop("Ooops.")
  value^2
}
batchMap(flakeyFunction, 1:10)</pre>
```

```
## Adding 10 jobs ...
```

Now that the jobs are defined, we can test jobs independently:

```
testJob(id = 1)
```

```
## [1] 1
```

In this case, testing the job with ID = 1 provides the appropriate result but testing the job with ID = 2 leads to an error:

```
as.character(try(testJob(id = 2)))
## [1] "Error in (function (value) : Ooops.\n"
We ignore the error here, and just assume everything looks fine and submit all jobs.
submitJobs()
## Submitting 10 jobs in 10 chunks using cluster functions 'Interactive' ...
## Warning in (function (value) : Just a simple warning
waitForJobs()
## [1] FALSE
After you have submitted jobs and suspect that something is going wrong, the first thing to do is to run
getStatus() to display a summary of the current state of the system.
getStatus()
## Status for 10 jobs:
     Submitted: 10 (100.0%)
##
##
              : 10 (100.0%)
     Started
##
     Done
                : 8 (80.0%)
                : 2 ( 20.0%)
##
     Error
##
     Queued
                : 0 ( 0.0%)
##
                   0 (
                        0.0%)
     Running
                   0 ( 0.0%)
     Expired
The status message shows that two of the jobs could not be executed successfully. To get the IDs of all jobs
that failed due to an error we can use findErrors() and to retrieve the actual error message, we can use
getErrorMessages().
findErrors()
##
      job.id
##
       <int>
## 1:
            9
## 2:
getErrorMessages()
##
      job.id terminated error
                                                                 message
```

```
## job.id terminated error message
## <int> <lgcl> <lgcl> < char>
## 1: 2 TRUE TRUE Error in (function (value) : Ooops.
## 2: 9 TRUE TRUE Error in (function (value) : Ooops.
```

If we want to peek into the R log file of a job to see more context for the error we can use **showLog()** which opens a pager or use **getLog()** to get the log as character vector:

```
tail(getLog(id = 9))
```

```
## [1] "### [bt 2017-09-06 15:16:19]: Memory measurement disabled"
## [2] "### [bt 2017-09-06 15:16:19]: Starting job [batchtools job.id=9]"
## [3] "Error in (function (value) : Ooops."
## [4] ""
## [5] "### [bt 2017-09-06 15:16:19]: Job terminated with an exception [batchtools job.id=9]"
## [6] "### [bt 2017-09-06 15:16:19]: Calculation finished!"
```

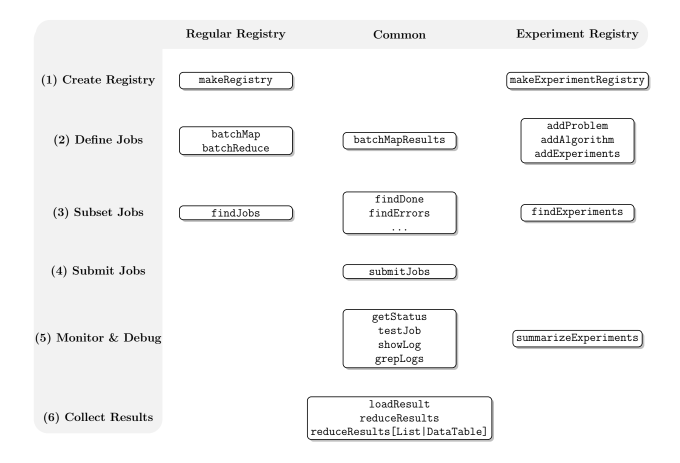
You can also grep for messages (output suppressed in this vignette for technical reasons):

```
grepLogs(pattern = "simple", ignore.case = TRUE)
```

Workflow

On the Local System

- 1. Create a Registry with makeRegistry() (or makeExperimentRegistry()) or load an existing from the file system with loadRegistry().
- 2. Define computational jobs with batchMap() or batchReduce() if you used makeRegistry() or define with addAlgorithm(), addProblem() and addExperiments() if you started with makeExperimentRegistry(). It is advised to test some jobs with testJob() in the interactive session and with testJob(external = TRUE) in a separate R process. Note that you can add additional jobs if you are using an ExperimentRegistry.
- 3. If required, query the data base for job ids depending on their status, parameters or tags (see findJobs()). The returned tables can easily be combined in a set-like fashion with data base verbs: union (ojoin() for outer join), intersect (ijoin() for inner join), difference (ajoin() for anti join).
- 4. Submit jobs with submitJobs(). You can specify job resources here. If you have thousands of fast terminating jobs, you want to chunk() them first. If some jobs already terminated, you can estimate the runtimes with estimateRuntimes() and chunk jobs into heterogeneous groups with lpt() and binpack().
- 5. Monitor jobs. getStatus() gives a summarizing overview. Use showLog() and grepLogs() to investigate log file. Run jobs in the currently running session with testJob() to get a traceback().
- 6. Collect (partial) results. loadResult() retrieves a single result from the file system. reduceResults() mimics Reduce() and allows to apply a function to many files in an iterative fashion. reduceResultsList() and reduceResultsDataTable() collect results into a list or data.table, respectively.



On Multiple Systems

Most users develop and prototype their experiments on a desktop box in their preferred IDE and later deploy to a large computing cluster. This can be done by prototyping locally (testJob() or submit subsets via submitJobs()). To deploy to the cluster, just copy the file directory (as reported by reg\$file.dir) to the remote system. Next, log in on the cluster (typically via ssh), cd to the copied directory and call loadRegistry("<file.dir.on.remote">, "<work.dir.on.remote>"). This function will (a) source the local configuration file so that you can talk to the cluster (verify by checking the output of reg\$cluster.functions) and (b) adjust the paths to the new system if argument update.paths is set. After loading the Registry, it is advised to test some jobs again with testJob() before submitting all of them with submitJobs(resources = list()) (remember you now need to set resources!). After some jobs are finished, the file.dir can be copied back (do not merge with the previous directory!) and loaded again with loadRegistry().

This approach is totally viable as long as some general rules are followed:

- 1. Make sure you have all packages installed. Package versions can be synchronized with checkpoint or packrat.
- 2. Test jobs on the remote system before submitting to ensure that paths are resolved correctly.
- 3. Make sure you have set the cluster functions in a configuration file, and stick to one backend as long as jobs are running.
- 4. The status can only be monitored on the remote system (for obvious reasons).
- 5. Partial results can be inspected both on the remote system and on the local system. For the latter, you need to copy over the **complete file.dir** first. Overwriting/merging directories is not advised as this may lead to inconsistencies if you added or removed experiments on the remote. If you have to merge, use rsync with option --delete. Load the registry locally with loadRegistry() and collect results.

Do not copy back and forth.

6. Never access the file.dir with multiple sessions simultaneously. Also never mount the file dir to work with it in a local R session. This may lead to inconsistencies and missing results.