How to use R on the Bioinformatics cluster
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DO NOT run treatments on frontal servers, always use sbatch or srun.

READ THE FAQ (http://bioinfo.genotoul.fr/index.php/faq/) before asking for support.

This tutorial aims at describing how to run R scripts and compile Rmarkdown files on the Toulouse Bioinformatics\(^1\) cluster. To do so, you need to have an account (ask for an account on this page http://bioinfo.genotoul.fr/index.php/ask-for/create-an-account/). You can then connect to the cluster using the ssh command on linux and Mac OS and using Putty\(^2\) on Windows. Similarly, you can copy files between the cluster and your computer using the scp command on linux and Mac OS and using WinSCP\(^3\) on Windows. The login address is genologin.toulouse.inra.fr. Once you are connected, you have two solutions to run a script: running it in batch mode or starting an interactive session. The script must never been run on the first server you connect to. Also, be careful that the programs that you can use from the cluster are not available until you have loaded the corresponding module. How to manage modules is explained in Section 1.

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1 Use of modules

All programs are made available by loading the corresponding module. These commands are the main useful commands to work with modules:

- module avail: list all available modules
- search_module [TEXT]: find a module with keyword

\(^1\)http://bioinfo.genotoul.fr/
\(^2\)https://putty.org/
\(^3\)https://winscp.net/eng/index.php
• **module load [MODULE_NAME]**: to load a module (for instance to load \texttt{R module load system/R-3.5.1}). This command is either used directly (in interactive mode) or included in the file that is used to run your \texttt{R} script in batch mode (see below)

• **module purge**: purge all previous loaded modules

2 **Run an \texttt{R} script in batch mode**

To launch an \texttt{R} script on the slurm cluster:

1. Write an \texttt{R} script:

   **Script 1: HelloWorld.R**

   ```
   print("Hello world!")
   ```

2. Write a \texttt{bash} script:

   **Script 2: myscript.sh**

   ```
   #!/bin/bash
   #SBATCH --job-name=launchRscript
   #SBATCH --output=output.out
   # Purge all previously loaded modules
   module purge
   # Load the \texttt{R} module
   module load system/R-3.5.1
   # The command lines that I want to run on the cluster
   Rscript HelloWorld.R
   ```

3. Launch the (bash) script with the \texttt{sbatch} command:

   ```
   sbatch myscript.sh
   ```

   The scripts \texttt{myscript.sh} and \texttt{HelloWorld.R} are supposed to be located in the same directory from which the \texttt{sbatch} command is launched. For \texttt{Rmd} files (see section 7), be careful that you cannot compile a document if the \texttt{Rmd} file is not in a writable directory.

2.1 **sbatch options**

Jobs can be launched with customized options (more memory, for instance). There are two ways to handle \texttt{sbatch} options:

• **[RECOMMENDED]** at the beginning of the \texttt{bash} script with lines of the form: \texttt{#SBATCH [OPTION] [VALUE]}

• in the \texttt{sbatch} command: \texttt{sbatch [OPTION] [VALUE] [OPTION] [VALUE] [...]} \texttt{myscript.sh}

Many options are available. To see all options use \texttt{sbatch --help}. Useful options:

• **-J, --job-name=jobname**: name of job

• **-e, --error=err**: file for batch script’s standard error

• **-o, --output=out**: file for batch script’s standard output

• **--mail-type=BEGIN,END,FAIL**: send an email at beginning, end or fail of the script (default email is your user email and can be changed with \texttt{--mail-user=truc@bidule.fr}, to use with care)
• -t, --time=HH:MM:SS: time limit (default to 04:00:00)
• --mem=XXG: to change memory reservation (default to 4G)
• -c, --cpus-per-task=ncpus: number of cpus required per task (default to 1)
• --mem-per-cpu=XXG: maximum amount of real memory per allocated cpu required by the job

2.2 Job management

After a job has been launched, you can monitor it with squeue -u [USERNAME] or squeue --j [JOB_ID] and also cancel it with scancel [JOB_ID].

3 Use R in an interactive mode

To use R in a console mode, use srun --pty bash to be connected to a node. Then, module load system/R-3.5.1 (for the last R version) and R to launch R.

srun can be run with the same options as for sbatch command (cpu and memory reservations) (see section 2.1).

3.1 X11 sessions

X11 sessions are useful to display plots directly in an interactive session. Prior their use, generate a ssh key with ssh-keygen and add it in authorized keys file with cat .ssh/id_rsa.pub >> .ssh/authorized keys. The interactive session is then launched with:

1. Logging on the cluster with ssh -Y [USERNAME]@genologin.toulouse.inra.fr
2. Run an interactive session with srun --x11 --pty bash
4 R in a parallel environment

To use R with a parallel environment, the `-c, --cpus-per-task=ncpus` option for the `sbatch` and `srun` is needed. In the R script the SAME number of cores needs to be specified.

4.1 Parallel with doParallel package

Two packages exist to use parallel with R: `doParallel` and `BiocParallel` (examples are provided for 2 parallel jobs).

1. Write an R script:

   • With `doParallel` package:
     
     ```R
     library(doParallel)
     cl <- makeCluster(2)
     registerDoParallel(cl)
     foreach(i=1:3) %dopar% sqrt(i)
     ```

   • With `BiocParallel` package:
     
     ```R
     library(BiocParallel)
     # specify the number of cores with workers = 2
     bplapply(1:10, print, BPPARAM = MulticoreParam(workers = 2))
     ```

2. Write a `bash` script:

   ```bash
   #!/bin/bash
   #SBATCH --job-name launchRscript
   #SBATCH --output output.out
   #SBATCH --cpus-per-task=2
   ```
#Purge any previous modules
module purge

#Load the application
module load system/R-3.5.1

# My command lines I want to run on the cluster
Rscript TestParallel.R

3. Launch the script with the `sbatch` command:
```bash
sbatch myscript.sh
```

## Arguments in a script

External arguments can be passed to an R script. The basic method is described below but the package `optparse` provides ways to handle external arguments à la Python.

1. Write an R script:

   **Script 6: HelloWorld.R**
   ```r
   args = commandArgs(trailingOnly=TRUE)
   print(args[1])
   ```

2. Write a `bash` script:

   **Script 7: myscript.sh**
   ```bash
   #!/bin/bash
   #SBATCH --J launchRscript
   #SBATCH --o output.out
   # Purge any previous modules
   module purge
   # Load the application
   module load system/R-3.5.1
   # My command lines I want to run on the cluster
   Rscript --vanilla HelloWorld.R "Hi!"
   ```

3. Launch the script with the `sbatch` command:
```bash
sbatch myscript.sh
```

## Install packages in your own environment

Once in an interactive R session, R packages are installed (in a personal library) using the standard `install.packages` command line.
Your personal library is usually located at the root of your personal directory whose allocated space is very limited. A simple solution consists in:

1. creating a directory named `R` elsewhere: `mkdir ~/work/R`
2. making a symbolic link to this directory: `ln -s ~/work/R ~/R`

7 Create and compile .Rmd (Rmarkdown) files on the cluster (batch mode)

To compile an .Rmd file, two packages are needed: `rmarkdown` and `knitr`. You also need to load the module `system/pandoc-2.1.3`.

As for an R script, you can pass external arguments to a .Rmd document.

1. Write an .Rmd script with parameters in the header:

   ```
   -----
   title : My Document
   output : html_document
   params:
   text : "Hi!"
   -----
   What is your text ?
   ```
   ```r
   print(params$text)
   ```

2. Write an R script to pass parameters:

   ```
   rmarkdown::render("MyDocument.Rmd", params = list(text = "Hola!"))
   ```

3. Write a bash script:
# Script 10: myscript.sh

```bash
#!/bin/bash
#SBATCH --J launchRscript
#SBATCH --o output.out

module purge
module load system/R-3.5.1
module load system/pandoc-2.1.3

Rscript --vanilla TestRmd.R
```

4. Launch the script with the `sbatch` command:

```bash
sbatch myscript.sh
```