

# **HPC** cluster Initiation

www.southgreen.fr

https://southgreenplatform.github.io/trainings















## i-Trop presentation













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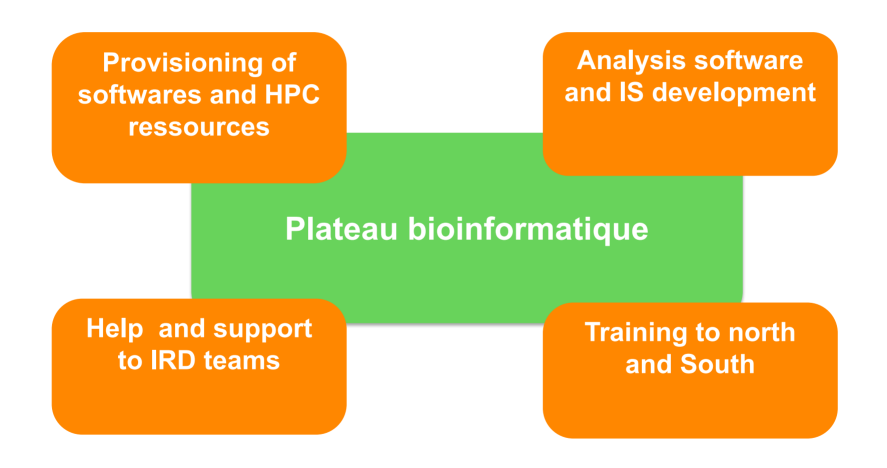
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## outh Green i-Trop Presentation





### **Demandes/incidents/Howtos**

### Request forms:

https://itrop-glpi.ird.fr/plugins/formcreator/front/formlist.php

- Accounts
- Softwares
- Projects



- Incidents: contact <u>bioinfo@ird.fr</u>
- Howtos:

https://southgreenplatform.github.io/trainings/hpc/hpcHowto/

### Slurm Tutorials:

https://southgreenplatform.github.io/tutorials//cluste
r-itrop/Slurm/



## **ARCHITECTURE**



### A Cluster?

- A logical unit composed of several servers
- A powerful unique machine
- Allow to obtain high computing performance
- A bigger capacity storage
- More reliable
- A better ressources availabilty



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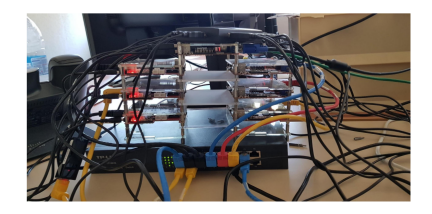




## A cluster?

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## **Cluster components**



- Master Node
   Handle resources and jobs
   priorities
- Computing nodes
   Resources (CPU or RAM memory)



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NAS Server(s)Storage



### 1 Master Node



bioinfo-master.ird.fr

### Role:

- Launch and prioritize jobs on computating nodes
- Accessible from the Internet
- Connection:

ssh login@bioinfo-master.ird.fr



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### 27 computing nodes



nodeX X: 0..26

### Role:

- Used by the master to execute jobs
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- node0 to node26



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### 27 computing nodes



nodeX X: 0..26

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### Interactif node (node6)

- Accessible from the Internet: bioinfo-inter.ird.fr
- Connection: ssh login@bioinfo-inter.ird.fr

# Practice

**Step 1: Connection, qhost** 

Go to the <a href="Practice1">Practice 1</a> of github



## Analyses steps of the cluster

Connection
to
bioinfo-mas
ter.ird.fr
and
resources
reservation

Step 1 salloc/srun ou sbatch

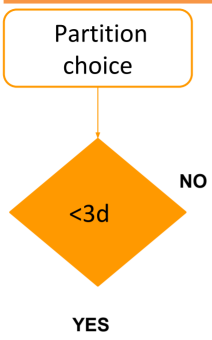


## **Partitions**

Partitions	Use	RAM on nodes	Core on nodes
short	Short Jobs < 1 day (higher priority, interactive jobs)	48 to 64 GB	12 cores
normal	Short Jobs max 3 days	64 Go to 96 GB	12 to 24 cores
long	45 days >long jobs > 3 days	48 GB	12 to 24 cores
highmem	Jobs with more memory needs	144 GB	12 to 24 cores
supermem	Jobs with much more memory needs	1TB	40 cores
gpu	Need for analyses on GPU cores	192GB	24 cpus and 8 GPUs cores

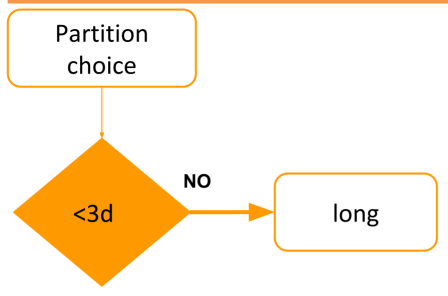


# outh Green How do I choose the partition?



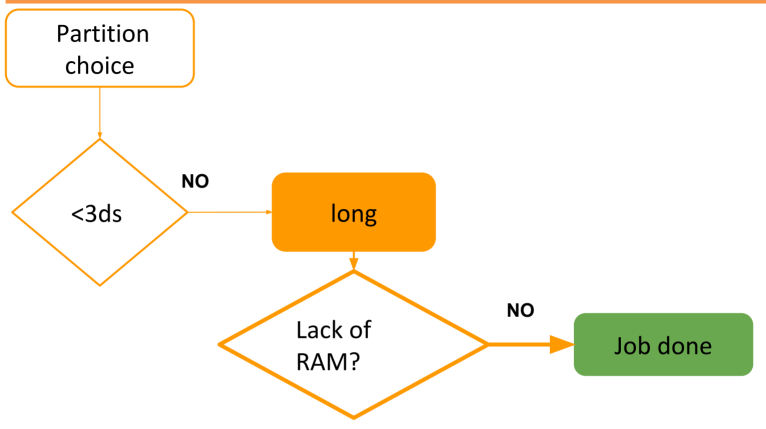


## How do I choose the partition?



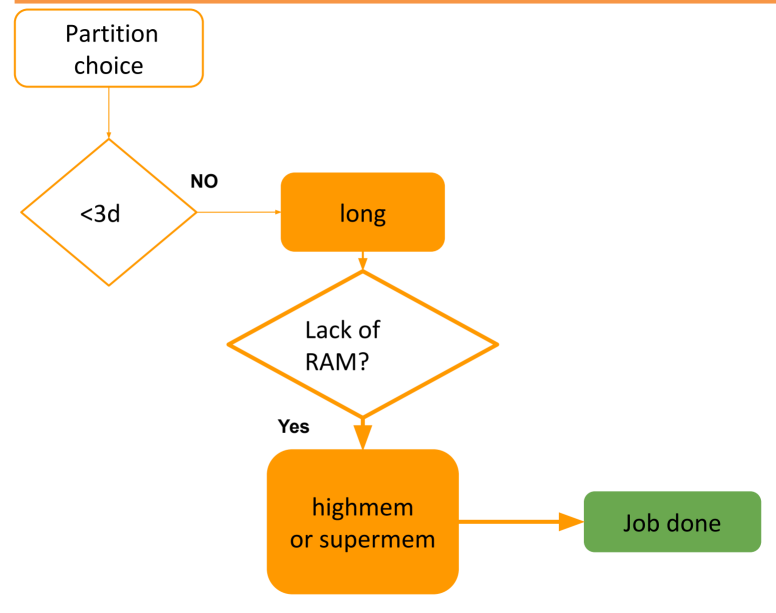


# outh Green How do I choose the partition?



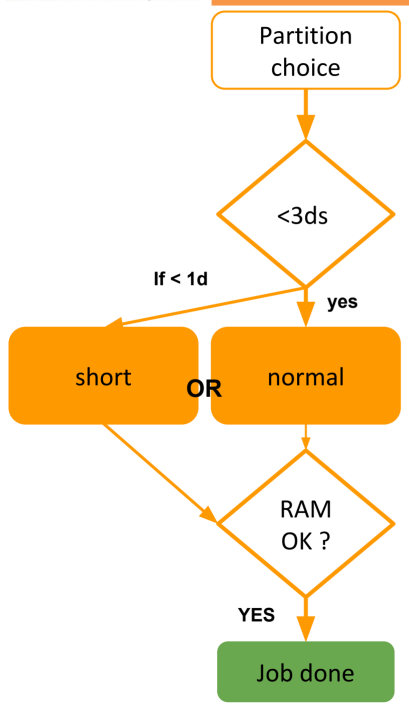


# outh Green How do I choose the partition?



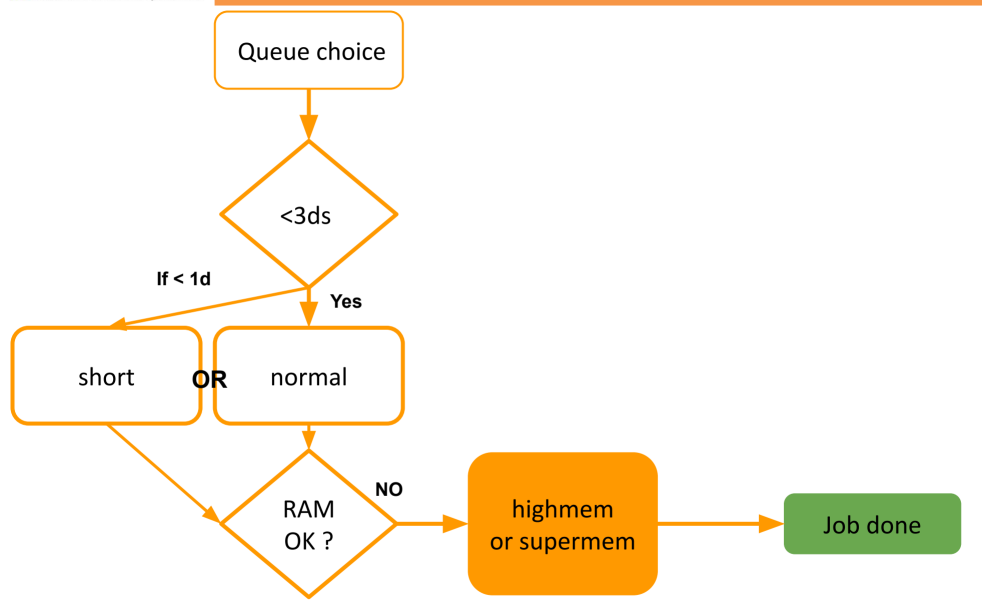


# uth Green How do I choose the partition?



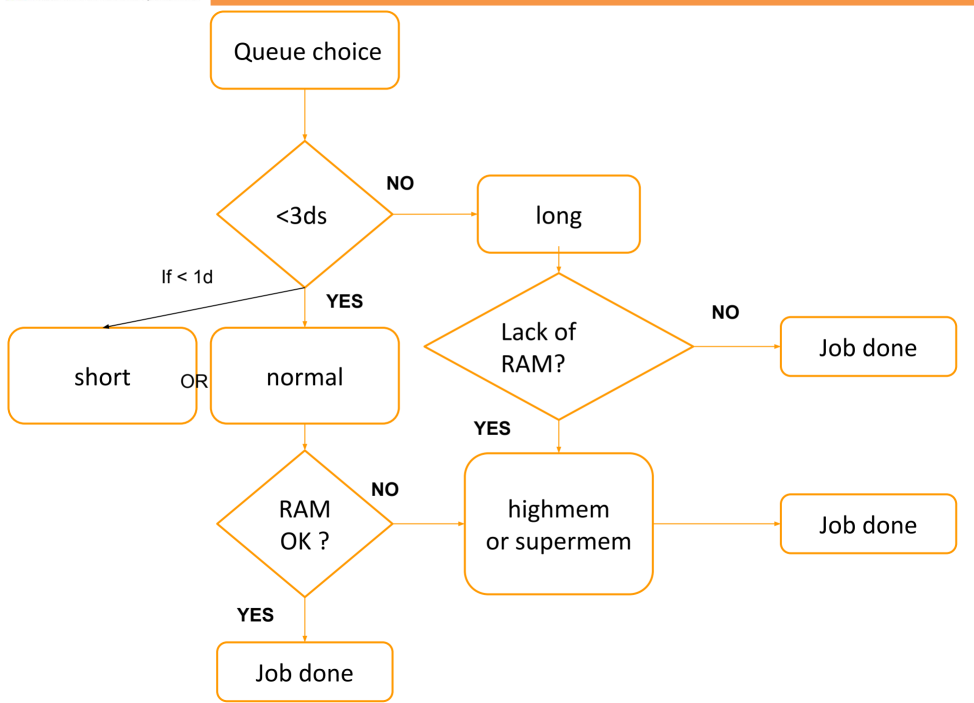


# outh Green How do I choose the queue?





## Green How do I choose the queue?





### 1 Master node



bioinfo-master.ird.fr

#### Role:

- Launch and prioritize jobs on computing nodes
- Accessible from the Internet

### **27** computing nodes



nodeX

X:0...26



#### Role:

- Used by the master to execute jobs
- Not accessible from the Internet

### 3 NAS servers



Bioinfo-nas.ird.fr (nas)

Bioinfo-nas2.ird.fr (nas2)

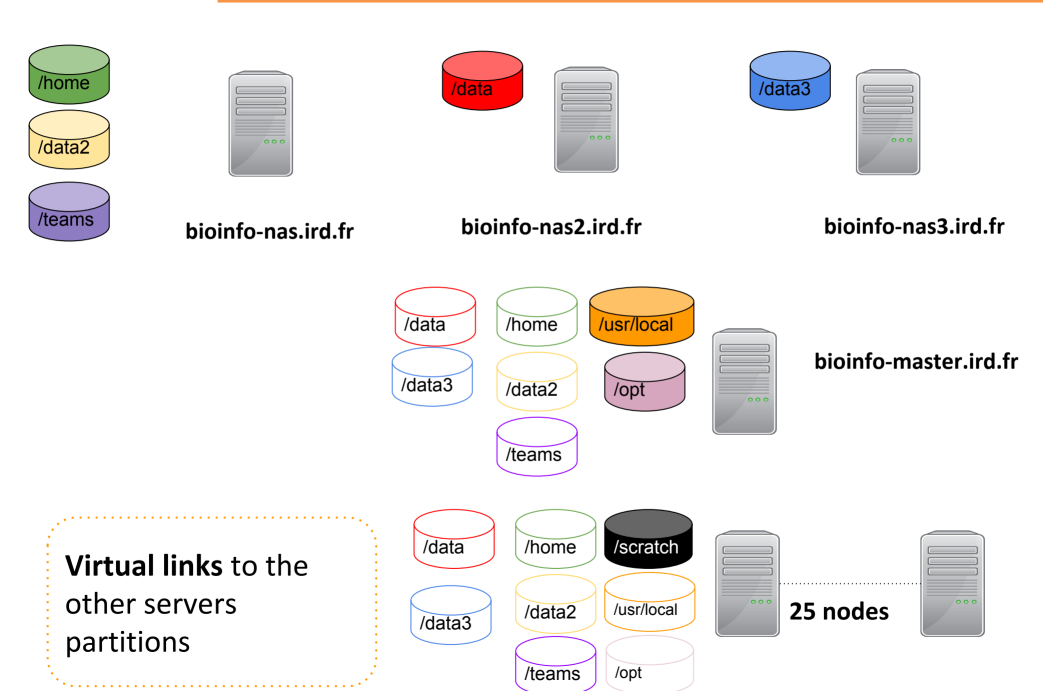
Bioinfo-nas3.ird.fr (nas3)

#### Role:

- Store users data
- Accessible from the Internet
- To transfer data: via filezilla or scp

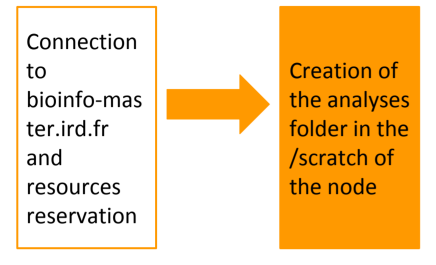


## cluster i-Trop disk partitions





## Analyses steps of the cluster



Step 1 Step 2 mkdir

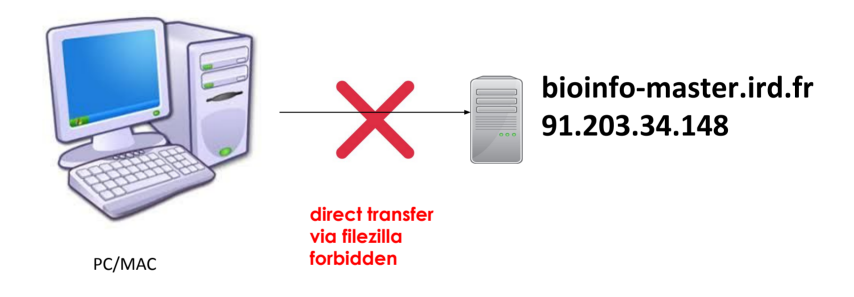
# Practice

Step 2:qrsh, partition

Go to the <a href="Practice2">Practice2</a> of the github



## Data transfer on i-Trop cluster





## Data transfer on i-Trop cluster

### /home and/or /teams or /data2

Hostname: bioinfo-nas.ird.fr

Login: cluster account

bioinfo-nas.ird.fr 91.203.34.157

Password: cluster

password Port : 22

/data



bioinfo-nas2.ird

91.203.34.160

.fr

Hostname: bioinfo-nas2.ird.fr

Login: cluster account

Password: cluster password

Port : 22

/data3



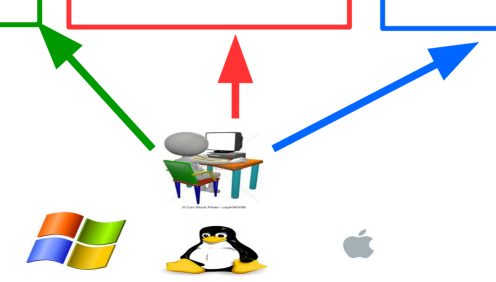
Hostname: bioinfo-nas3.ird.fr

Login: cluster account

91.203.34.180

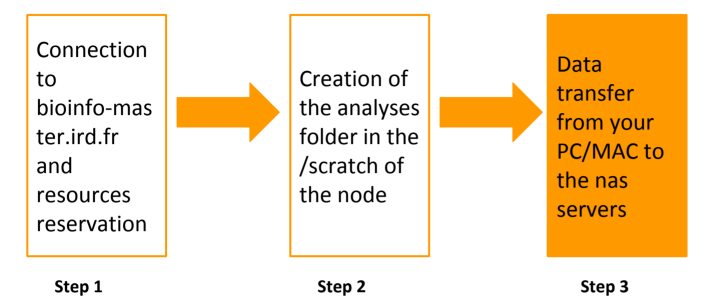
Password: bioinfo-nas3.ird.cluster password

Port : 22





## Analyses steps of the cluster





Copy your data from your PC/MAC to the nas serve if they are not on the cluster

# Practice

Step 3: filezilla

Go to the <a href="Practice3">Practice3</a> of the github

## Copy with scp

Copy between 2 remote servers :

scp -r source destination

• Syntax if the source is remote:

scp -r server\_name:/path/file\_to\_copy local\_folder

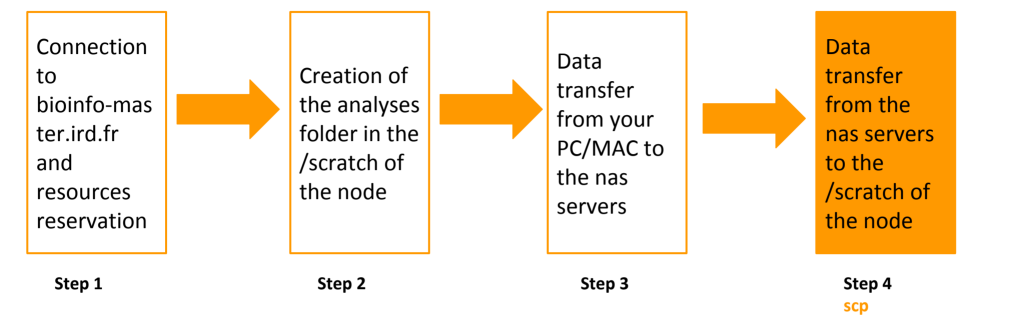
Syntax if the destination is remote :

scp -r /path/file\_to\_copy server\_name:/path/remote\_folder

Ex: scp -r nas:/home/tando/folder//scratch/tando/



## **Analyses steps of the cluster**



# Practice

**Step 4: scp to nodes** 

Go to the <a href="Practice4">Practice4</a> of the github



### **Module Environment**

- Allow to choose the version of software you want to use
- 2 types of softwares: bioinfo: includes all the bioinformatics softwares ( example BEAST) system: includes all the system softwares(example JAVA)
- Overcome the environment variables



### **Module Environment**

- 5 types of commands :
- See the available modules :

module avail

Obtain infos on a particular module:
 module whatis + module name

Load a module :

module load + modulename

List the loaded module :

module list

• Unload a module:

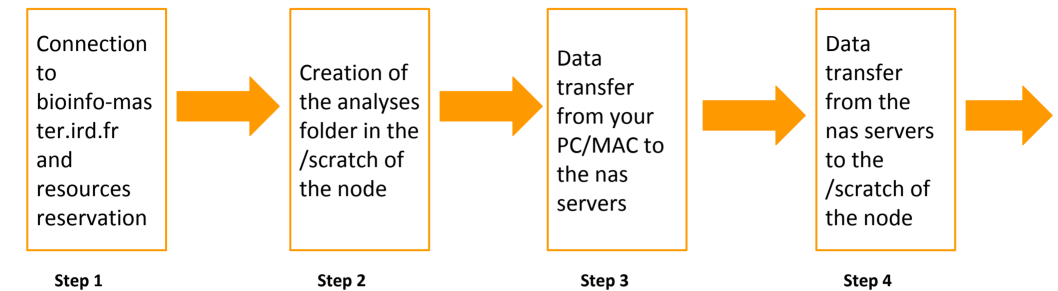
module unload + modulename

Unload all the modules :

Module purge



### Analyses steps of the cluster



Load softwares with modules environment

Step 5 module

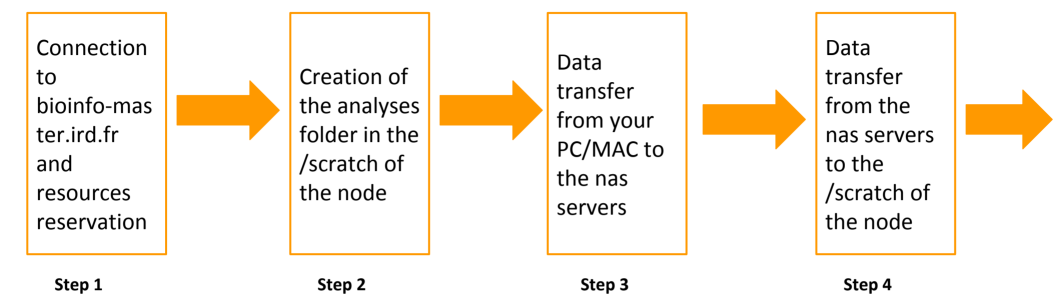
## **Practice**

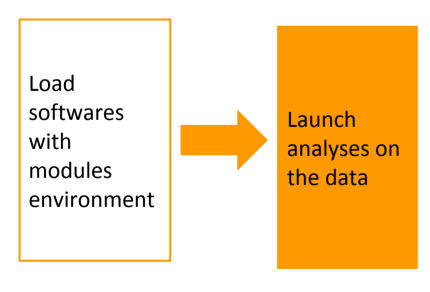
**Step 5: module environment** 

Go to the <a href="Practice5">Practice5</a> of the github



### Analyses steps of the cluster





Step 5 Step 6

## South Green Main Slurm commands Main Slurm commands

Commande	Description	Exemple
sruntime=0X:00pty bash -i	Interactive way to connect to a node for X minutes	sruntime=02:00:00pty bash -i Connection for 2h
salloctime=0X:00	Allocate one or several nodes for a later use	Salloc -N 2p shorttime=05:00
sbatch	Launch a analyses in backkground via a script	sbatch script.sh
sinfo	Informations on partitions	sinfo
sinfo -N I	Informations on nodes of the partitions	sinfo -N l
squeue	Infos on all jobs	squeue -u tando
scontrol show job <job_id></job_id>	Infos on the active job <job_id></job_id>	scontrol show job 1029



## South Green Options des commandes sbatch, srun, salloc

Options	Description	Exemple
job-name= <name></name>	Name the job	sbatchjob-name=tando_blast
-p <partition></partition>	Choose a partition	sbatch -p highmem
nodelist= <nodex></nodex>	Choisir a particular node	Sbatch -p normalnodelist=node14
-n <nbre_cpus></nbre_cpus>	Launch with several cores	srun -n 4
mail-user= <emailaddress></emailaddress>	Send a email	sbatch mail-user=ndomassi.tando@ird.f r
mail-type= <event></event>	Send a email when: END: end of the job FAIL: abortion BEGIN: beginning of job	Sbatchmail-type=BEGIN
workdir=[dir_name]	Precise the working directory	sbatch sworkdir=/scratch/tando script.sh

Plus d'infos sur Slurm ici: <a href="https://southgreenplatform.github.io/tutorials//cluster-itrop/Slurm/#part-2">https://southgreenplatform.github.io/tutorials//cluster-itrop/Slurm/#part-2</a>



### Launch a command from the prompt

- Load the software version to launch
- Launch the data analysis

\$~ command <options> <arguments>

With command: the command to launch



### Launch a command line job

- Execute a bash command via qsub
- Launch the command from a node
- We use:

With command: the command to launch

## **Practice**

**Step6:** launch the analysis

Go to the <a href="Practice6">Practice6</a> of the github

### Transfer your results to the nas servers

Copy between 2 remote servers :

scp source destination

Syntax if the source is remote :

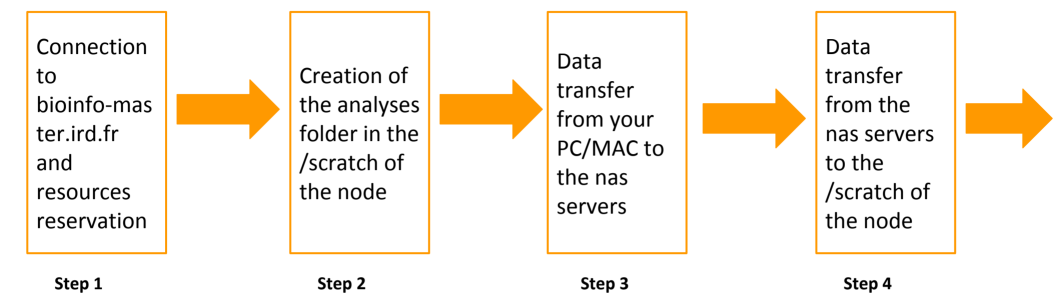
scp server\_name:/path/file\_to\_copy local\_folder

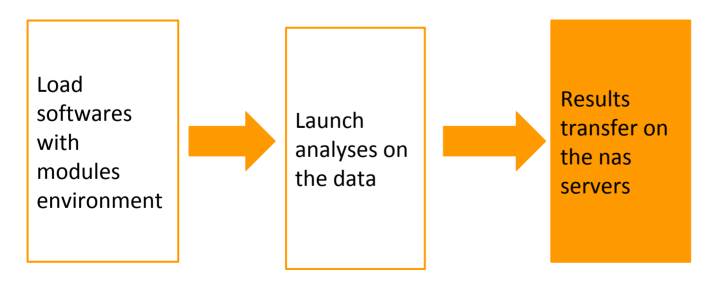
• Syntax if the destination is remote:

scp /path/file\_to\_copy server\_name:/path/remote\_folder



### Analyses steps of the cluster





Step 5 Step 6 Step 7 scp

## **Practice**

### **Step 7: Retrieve the results**

Go to the <a href="Practice7">Practice7</a> of the github



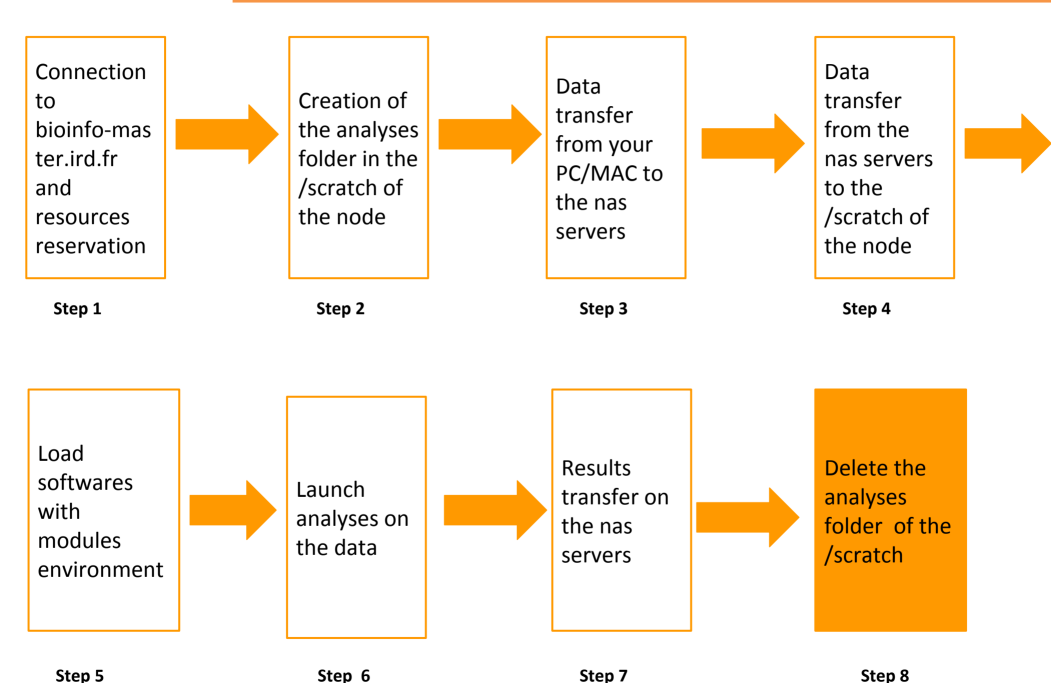
### **Delete results from scratchs**

- Scratch= temporary spaces
- Verify that the copy is OK before
- Use rm command

```
cd /scratch
rm -rf nom_rep
```



### Analyses steps of the cluster



rm

## **Practice**

**Step8: Data deletion** 

Go to the <a href="Practice8">Practice8</a> of the github



## Scripts to visualize/delete données temporary data

- Scripts location: /opt/scripts/scratch-scripts
- Visualize data on scratchs: scratch\_use.sh

sh /opt/scripts/scratch-scripts/scratch\_use.sh

Delete data on scratchs: clean\_scratch.sh

sh /opt/scripts/scratch-scripts/clean\_scratch.sh



## **BONUS**



### **LAUNCH A JOB**



### Advantages

- Scheduler choose resources automatically
- Use up to 24 cores at the same time
- Possibility to configure this choice
- Jobs launch in background
  - → possibility to turn off your PC/MAC
  - → automatic results retrieving



### Launch a batch job

- Execute a script via rm
- Use:

\$~ sbatch script.sh

with script.sh: the name of the script



## South Green Options des commandes sbatch, srun, salloc

Options	Description	Exemple
job-name= <name></name>	Name the job	sbatchjob-name=tando_blast
-p <partition></partition>	Choose a partition	sbatch -p highmem
nodelist= <nodex></nodex>	Choisir a particular node	Sbatch -p normalnodelist=node14
-n <nbre_cpus></nbre_cpus>	Launch with several cores	srun -n 4
mail-user= <emailaddress></emailaddress>	Send a email	sbatch mail-user=ndomassi.tando@ird.f r
mail-type= <event></event>	Send a email when: END: end of the job FAIL: abortion BEGIN: beginning of job	Sbatchmail-type=BEGIN
workdir=[dir_name]	Precise the working directory	sbatch sworkdir=/scratch/tando script.sh

Plus d'infos sur Slurm ici: <a href="https://southgreenplatform.github.io/tutorials//cluster-itrop/Slurm/#part-2">https://southgreenplatform.github.io/tutorials//cluster-itrop/Slurm/#part-2</a>



#!/bin/bash

### **Bash scripts syntax**

First part of the script (in green): sge execution options with the key word #SBATCH



### **Bash scripts syntax**

#### In the 2nd part of the script: the command to execute

nom variable1="valeur variable1" nom variable2="valeur variable2" sleep 30 hostname

## **Practice**

### Launch a script with sge

Go to the <a href="Practice9">Practice9</a> of the github

### Citations

If you use i-Trop Bioinformatics resources.

Thank you for citing with:

"The authors acknowledge the IRD itrop HPC (South Green Platform) at IRD montpellier

for providing HPC resources that have contributed to the research results reported within this paper.

URL: https://bioinfo.ird.fr/- http://www.southgreen.fr"

### **Projects**

Include a budget for bioinformatics resources in your answer to projects funding

- A need in hard drives, renewal machinesetc...
- Available quotations

 Contact <u>bioinfo@ird.fr</u>: help, needs definition, quotations...



# Thank you for your attention!



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