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## Formation FROGS juillet 2019

Géraldine Pascal, Laurent Cauquil

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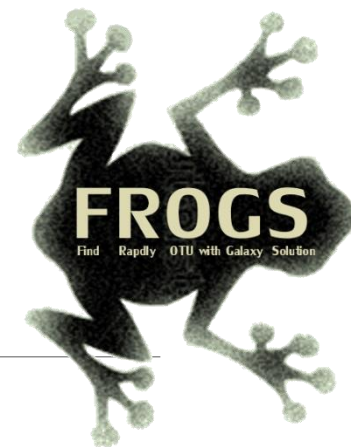
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**<https://hal.inrae.fr/hal-02790029>**

Submitted on 5 Jun 2020

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# Training on Galaxy: Metagenomics

July 2019

## Find, Rapidly, OTUs with Galaxy Solution

FRÉDÉRIC ESCUDIÉ\* and LUCAS AUER\*, MARIA BERNARD, LAURENT CAUQUIL, SARAH MAMAN, MAHENDRA MARIADASSOU, SYLVIE COMBES, GUILLERMINA HERNANDEZ-RAQUET, GÉRALDINE PASCAL & MALO LE BOULCH

\*THESE AUTHORS HAVE CONTRIBUTED EQUALLY TO THE PRESENT WORK.



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# Introduction to Galaxy platform and preparation of FROGS training

July 2019

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MALO LE BOULCH, SARAH MAMAN, GÉRALDINE PASCAL

# Chitchat time!

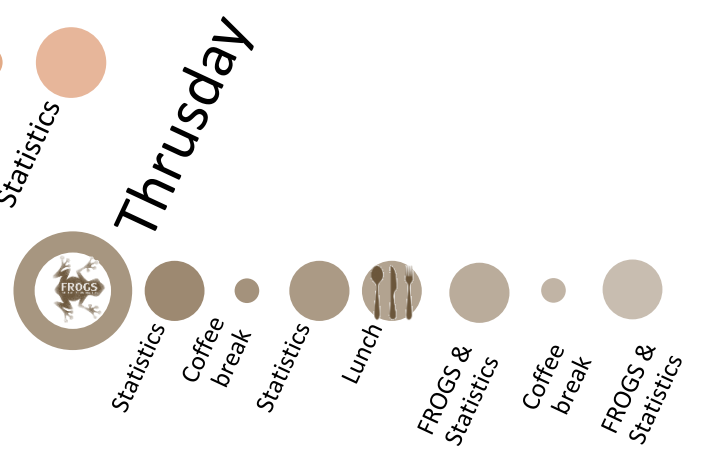
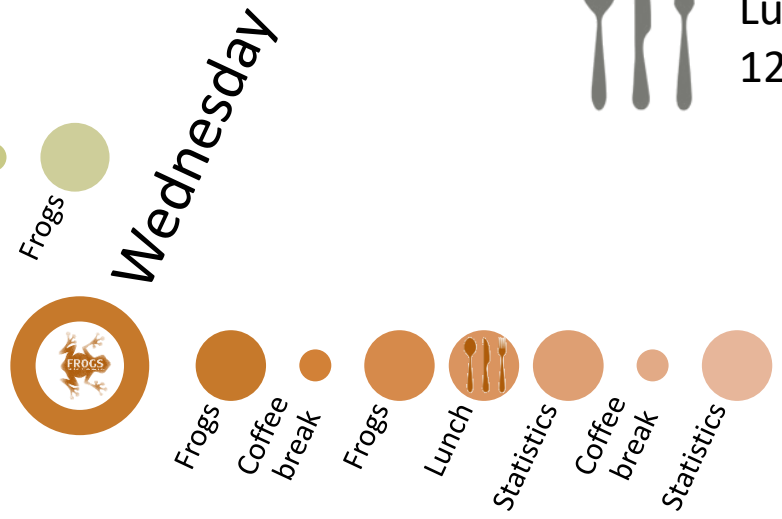
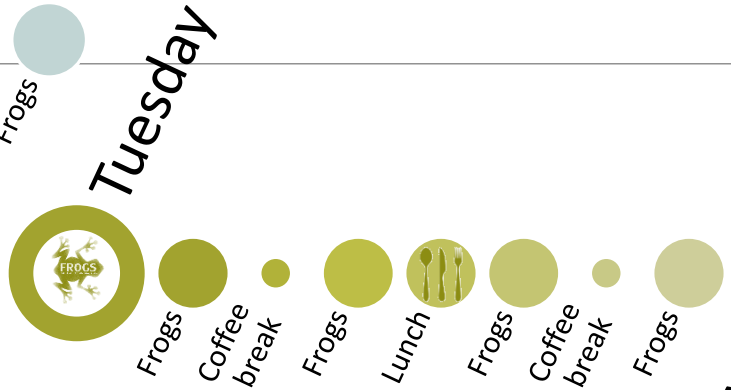
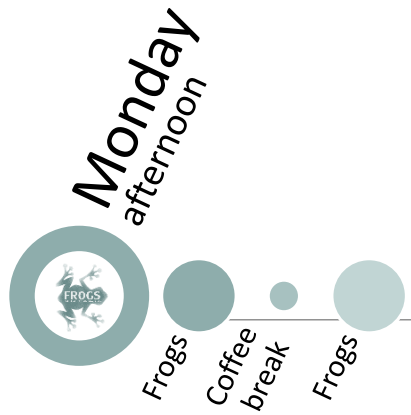
---

- What is your computer skills level?
- Have you ever heard of or used Galaxy?

# Feedback on metagenomics:

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- What are your needs in “metagenomics”?
- 454 / MiSeq?
- 16S/ITS ?
- Your background ?



9 am to 5 pm



2 short coffee breaks  
morning and afternoon



Lunch  
12.30 to 2.00 pm

# Objectives of the 1<sup>st</sup> part

---

- Learn the basics of Galaxy
- Being independent when using it
- Prepare the datasets for FROGS formation

# What is Galaxy?

---

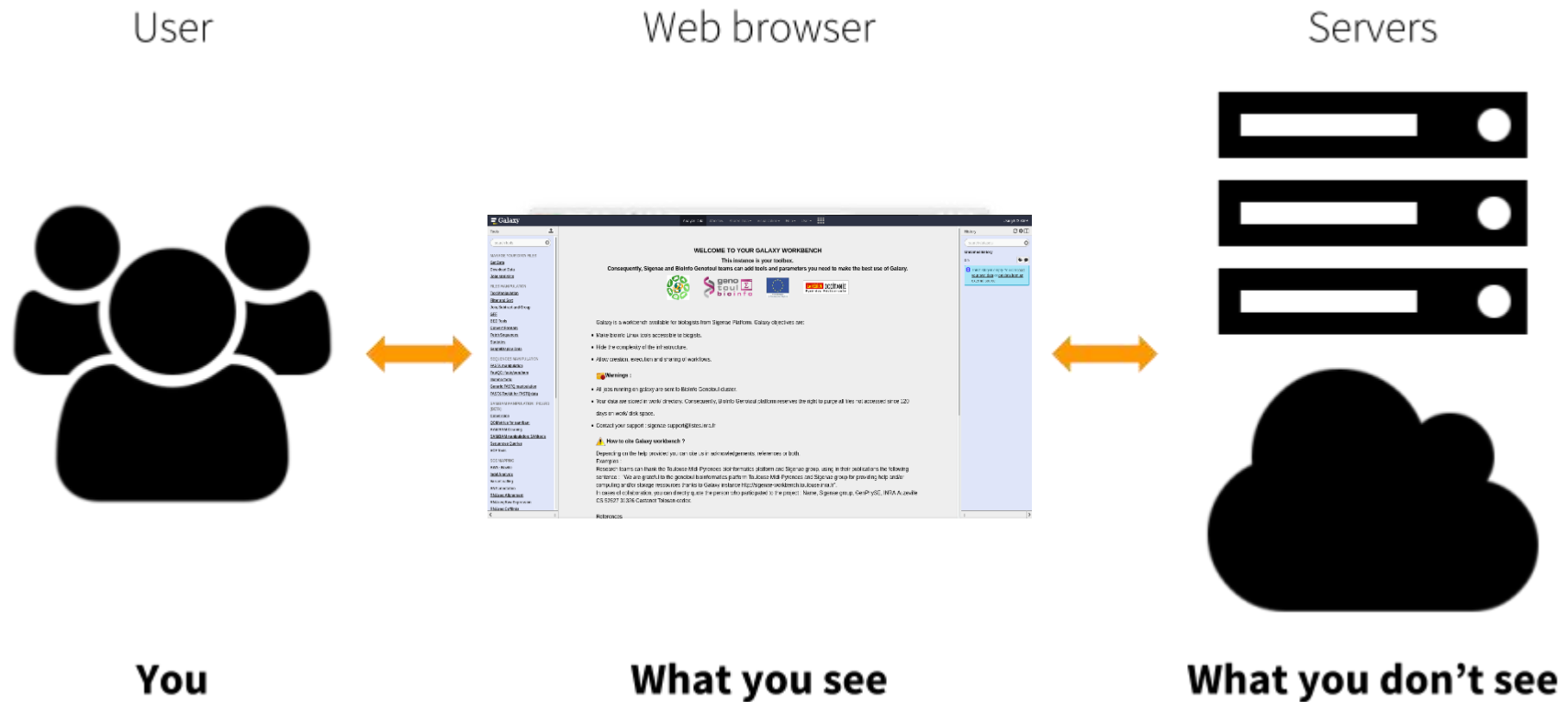


# Galaxy software framework

---

- Galaxy is an overlay that allows researchers without computer expertise to perform computational analyses online, automate them and share the results easily.
- Developed by the Galaxy Team Project.
- Relies on a computational infrastructure (Server).
- Open source

# Galaxy software framework



<https://galaxyproject.org/tutorials/g101/>

# Where to use Galaxy?

---

- Galaxy software can be installed and run on powerful server farms (Cluster) or on your individual PC.

- Genotoul Bioinfo



- Cluster: Many computers tightly connected that work together
- High performance computer:
  - More than 5000 cores
  - 34 TB of RAM
  - More than 1 Peta Byte (1024 TB) of hard drive



# Where to use Galaxy?

---

- Our Galaxy platform is <http://sigenae-workbench.toulouse.inra.fr/galaxy/>
- ⚠ The different platforms of Galaxy are not connected together
  - Galaxy is installed on many clusters across the world.
  - Some tools are in our Galaxy platform but not in other platforms.
  - Your data is not shared with other Galaxy platforms than ours.

# Exemple of 2 INRA Galaxy platforms

**Galaxy / Migale** Analyze Data Workflow Shared Data Visualization Help User

Tools

search tools

**Migale Tools**

- Get Data
- Send Data
- Lift-Over
- Gene Annotation
- Gene Prediction
- Text Manipulation
- Filter and Sort
- Join, Subtract and Group
- GFFtools
- Convert Formats
- Extract Features
- Fetch Sequences
- Fetch Alignments
- Get Genomic Scores
- Statistics
- Graph/Display Data
- Phenotype Association
- Sequence Alignment/Pairwise Alignment
- Metagenomic analyses
  - Metagenomics FROGS
  - Metagenomics Qiime
  - Metagenomics Mothur
- FASTA manipulation
- NGS: QC and manipulation
- NGS: Assembly
- NGS: Mapping
- NGS: RNA Analysis
- NGS: SAM Tools
- NGS: Peak Calling
- SNP/WGA: Data: Filters
- Variant Analysis
- NGS: Picard
- NGS: Variant analysis

**Workflows**

- All workflows

✓ Bienvenue sur le portail Galaxy de la plateforme Migale. Pour tous renseignements, demandes ou remarques, veuillez contacter [galaxy-help@jouy.inra.fr](mailto:galaxy-help@jouy.inra.fr)

**migale**  
Plateforme de BioInformatique - INRA Jouy en Josas

**Galaxy** est une plateforme qui propose une « constellation » d'outils pour analyser, manipuler et visualiser des données génomiques, sans avoir besoin de connaissance en programmation. Elle est développée par The Center for Comparative Genomics and Bioinformatics. L'utilisateur peut réaliser quatre grands types d'opérations :

- **manipulation de fichiers** : ajout ou suppression de colonnes, trier les fichiers, concaténer plusieurs fichiers, comparaison de listes, ...
- **opérations sur les données** : sommer, moyenner, soustraire, calculer la couverture d'une région déterminée, ...
- **analyse de séquences** : calculer des corrélations, utiliser des outils d'EMBOSS, aligner les données de séquençage, ...
- **visualisation des données** : afficher des alignements multiples, générer des graphiques, ...

**Contact** : [galaxy-help@jouy.inra.fr](mailto:galaxy-help@jouy.inra.fr)

Pour toutes demandes d'intégration de nouvel outil au sein du portail, veuillez remplir le [formulaire](#) mis à votre disposition sur le [site web de la plateforme Migale](#).

Une [Foire Aux Questions](#) autour de Galaxy a été mise en place sur le [site web de la plateforme Migale](#).

Si vous utilisez le portail Galaxy de la plateforme Migale pour effectuer vos analyses, merci de [citer Galaxy](#) ET de [remercier la plateforme dans les acknowledgements](#).

INRA  
SCIENCE & IMPACT

This project is supported in part by NSF, NHGRI, and the Huck Institutes of the Life Sciences.

# Exemple of 2 INRA Galaxy platforms

**Galaxy** | Analyze Data | Workflow | Shared Data | Visualization | Help | User | Using 0%

Tools | search tools

MANAGE YOUR DATA FILES  
[Get Data](#)  
[Download Data](#)  
[Jobs statistics](#)

FILES MANIPULATION  
[Text Manipulation](#)  
[Filter and Sort](#)  
[Join, Subtract and Group](#)  
[GFF](#)  
[BED Tools](#)  
[Convert Formats](#)  
[Fetch Sequences](#)  
[Statistics](#)  
[Graph/Display Data](#)

SEQUENCES MANIPULATION  
[FASTA manipulation](#)  
[FastQC: fastq/sam/bam](#)  
[Illumina fastq](#)  
[Generic FASTQ manipulation](#)  
[FASTX-Toolkit for FASTQ data](#)

SAMBAM MANIPULATION : PICARD (BETA)  
[Conversion](#)  
[QC/Metrics for sam/bam](#)  
[BAM/SAM Cleaning](#)  
[SAMBAM manipulation: SAMtools](#)  
[Sequences Queries](#)  
[VCF Tools](#)

SGS MAPPING  
[BWA - Bowtie](#)  
[Indel Analysis](#)  
[Variant calling](#)  
[SNP annotation](#)  
[RNAseq Alignment](#)  
[RNAseq Raw Expression](#)  
[RNAseq Cufflinks](#)

Genotoul Bioinfo | European Union | LA REGION OCCITANIE Pyrénées-Méditerranée

Welcome on Galaxy Sigene / BioInfo Genotoul.  
Contact your support : support.sigene@inra.fr

**Ready to use Workflows**

- 1/ FROGS - Find, Rapidly, Otus with Galaxy Solution
  - 2/ Tax4Fun workflow
- 3/ Function Table for Tax4Fun matrix workflow

**Galaxy News**

- 1/ How to use datasets collections ?
- 2/ How to upload several files in Galaxy ?
- 3/ New tools : Tax4Fun (manual), VIP (manual), Salmon, multiQC.
- 4/ Tools availables for virologie analyses
  - 5/ Virology tools
- 6/ How to use Function Table tool ?
- 7/ How to export and import your histories ?

**E-learning**

<https://inra.classilio.com/Login>  
2 e-learning availables : Galaxy initiation and Quality sequences analysis

Training resources : [BioInfo Genotoul / Sigene](#)

History | search datasets | 33 b

Data Collection | 3 shown

- 3: seq1.fasta
- 2: seq2.fasta
- 1: seq3.fasta

# Your Turn!

---

CONNECT TO OUR GALAXY WORKBENCH

# Exercise

---

During this formation, we will use a remote computer.

Connect to a distant computer by clicking on the remote desktop connection icon in the task bar.



Ask trainers for the address and password.

**All your data will be  
erased at the end of the  
week of the remote  
computer**



# Exercise


---

During this formation if you have a personal account, use it.

This account will save all the work done during this week and your future work.

If you don't have a personal account ask us for a temporary account.

You can request a personal account later on : <http://bioinfo.genotoul.fr/index.php/ask-for/create-an-account/>

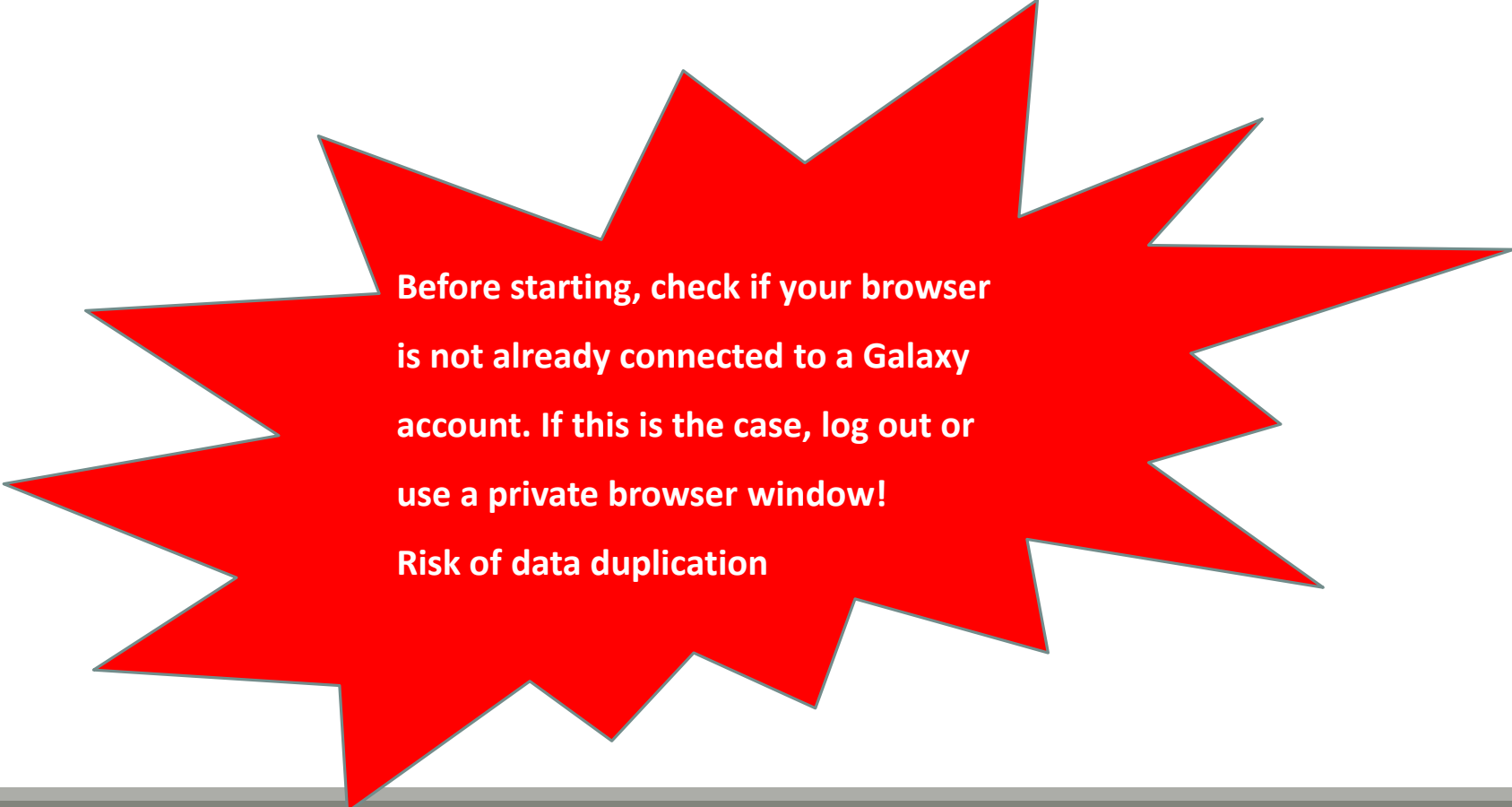


**All your data will be  
erased at the end of the  
week on a temporary  
account**

# Exercise

---

Our Galaxy platform is: <http://sigenae-workbench.toulouse.inra.fr/galaxy/>

A large, red, multi-pointed starburst graphic with a black outline, centered on the slide. It contains white text.

**Before starting, check if your browser  
is not already connected to a Galaxy  
account. If this is the case, log out or  
use a private browser window!  
Risk of data duplication**

# Exercise

---

Our Galaxy platform is: <http://sigenae-workbench.toulouse.inra.fr/galaxy/>

Be careful, to fully login you must enter your credentials twice:

- The first time in this pop-up window:



Authentication requise

 Le site <http://galaxy-workbench.toulouse.inra.fr> demande un nom d'utilisateur et un mot de passe. Le site indique : « Please enter your Genotoul LDAP password »

Utilisateur :

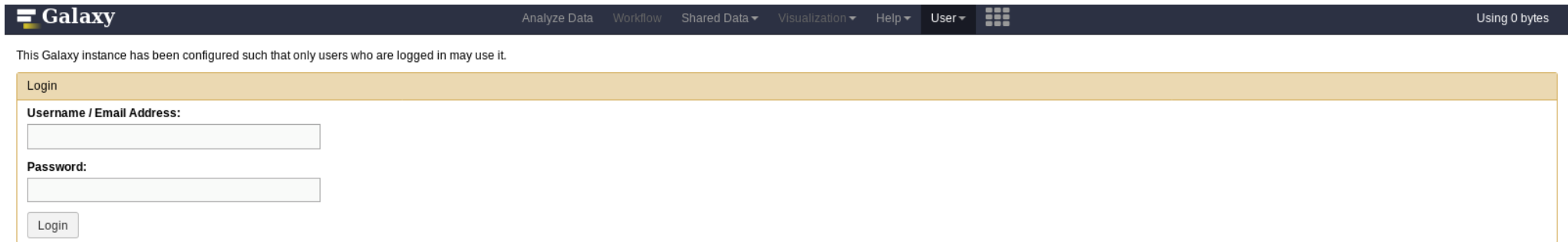
Mot de passe :

Annuler OK

# Exercise

---

- And a second time, in the browser:



The screenshot shows the Galaxy web interface. At the top, there is a dark navigation bar with the Galaxy logo on the left and several menu items: 'Analyze Data', 'Workflow', 'Shared Data', 'Visualization', 'Help', and 'User'. On the far right of the navigation bar, it says 'Using 0 bytes'. Below the navigation bar, a message reads: 'This Galaxy instance has been configured such that only users who are logged in may use it.' Below this message is a light-colored box containing a login form. The form has a title 'Login' in the top left corner. It contains two input fields: the first is labeled 'Username / Email Address:' and the second is labeled 'Password:'. Below the password field is a 'Login' button.

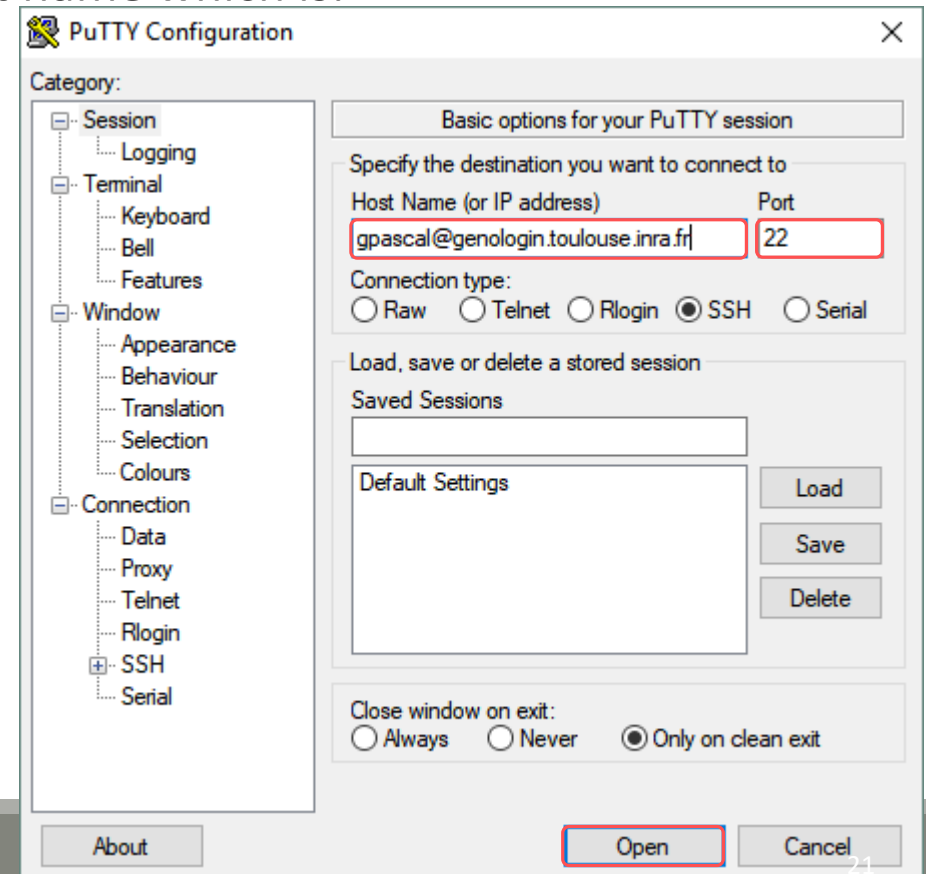
# Change your password

---

- In order to change your password, you need to download PuTTY: <http://www.putty.org/>
- PuTTY is a terminal emulator, it allows to connect directly to the server in command line.
- You can not change your password via the Galaxy's interface for the moment.

# Change your password

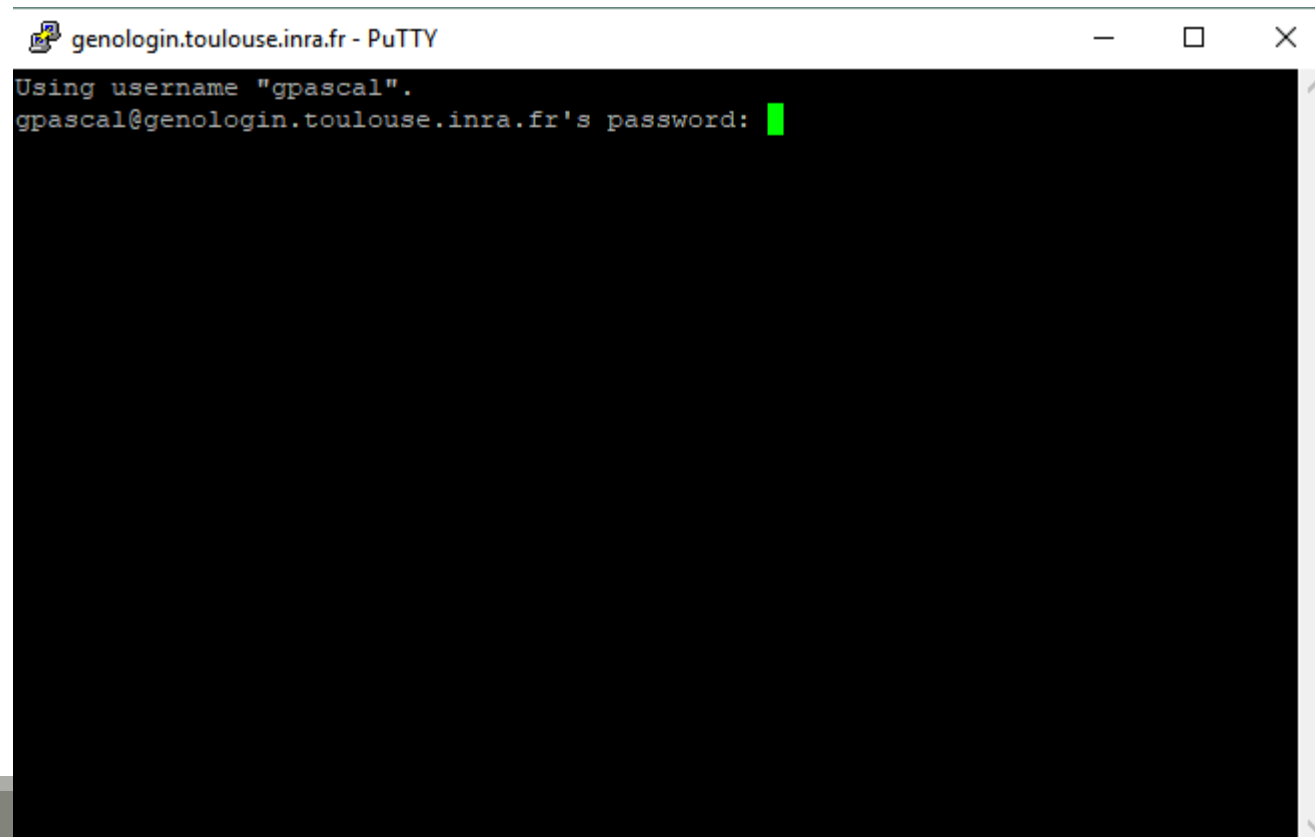
- Launch PuTTY.
- In the following window, you must enter your host name which is:  
[YourGenotoulName@genologin.toulouse.inra.fr](mailto:YourGenotoulName@genologin.toulouse.inra.fr)
- The connection type must be SSH.
- Click on « Open » .



# Change your password

---

- A new window appear, click on « Yes ».
- In the following window, enter your password and hit « Enter ».



The screenshot shows a PuTTY terminal window titled "genologin.toulouse.inra.fr - PuTTY". The terminal output is as follows:

```
Using username "gpascal".  
gpascal@genologin.toulouse.inra.fr's password: █
```

The terminal background is black, and the text is white. A green cursor is visible at the end of the password prompt line.

# Change your password

- Type « passwd » and hit « Enter ».



```
genologin.toulouse.inra.fr - PuTTY
e cluster).
Pour obtenir plus de memoire, veuillez consulter la FAQ de notre site web (cf +
bas)

=====
Informations concernant le quota de temps de calcul
=====

Il existe un quota de temps de calcul annuel de 100.000H pour les academiques (5
00H pour les entreprises privees).
Au dela il faut renseigner le formulaire de demande de ressources exceptionnelle
s sur notre site web.

Vous pouvez verifier votre quota de calcul avec la commande: squota_cpu

=====
Support
=====

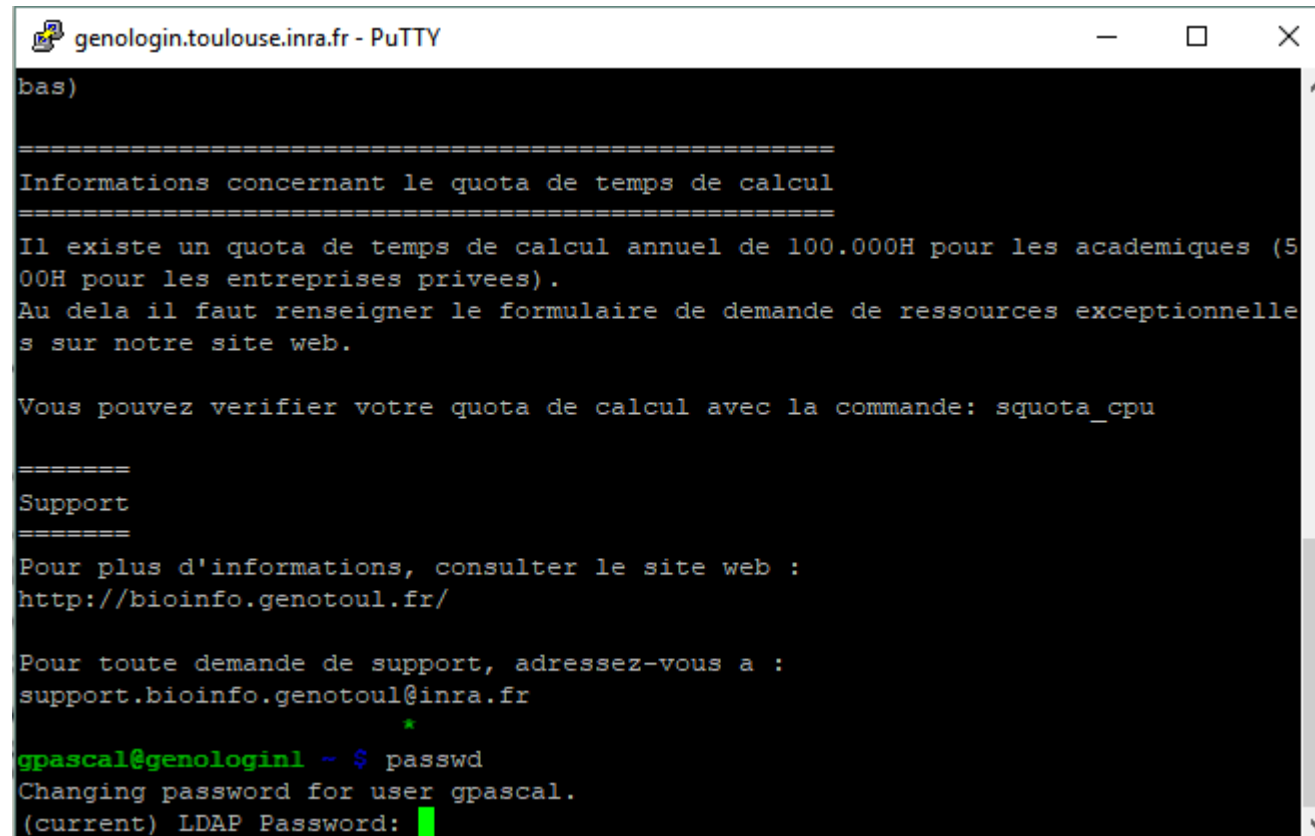
Pour plus d'informations, consulter le site web :
http://bioinfo.genotoul.fr/

Pour toute demande de support, adressez-vous a :
support.bioinfo.genotoul@inra.fr
*
gpascal@genologin1 ~ $ passwd
```



# Change your password

- Enter your current password and hit « Enter ».



```
genologin.toulouse.inra.fr - PuTTY
bas)

=====
Informations concernant le quota de temps de calcul
=====

Il existe un quota de temps de calcul annuel de 100.000H pour les academiques (5
00H pour les entreprises privees).
Au dela il faut renseigner le formulaire de demande de ressources exceptionnelle
s sur notre site web.

Vous pouvez verifier votre quota de calcul avec la commande: squota_cpu

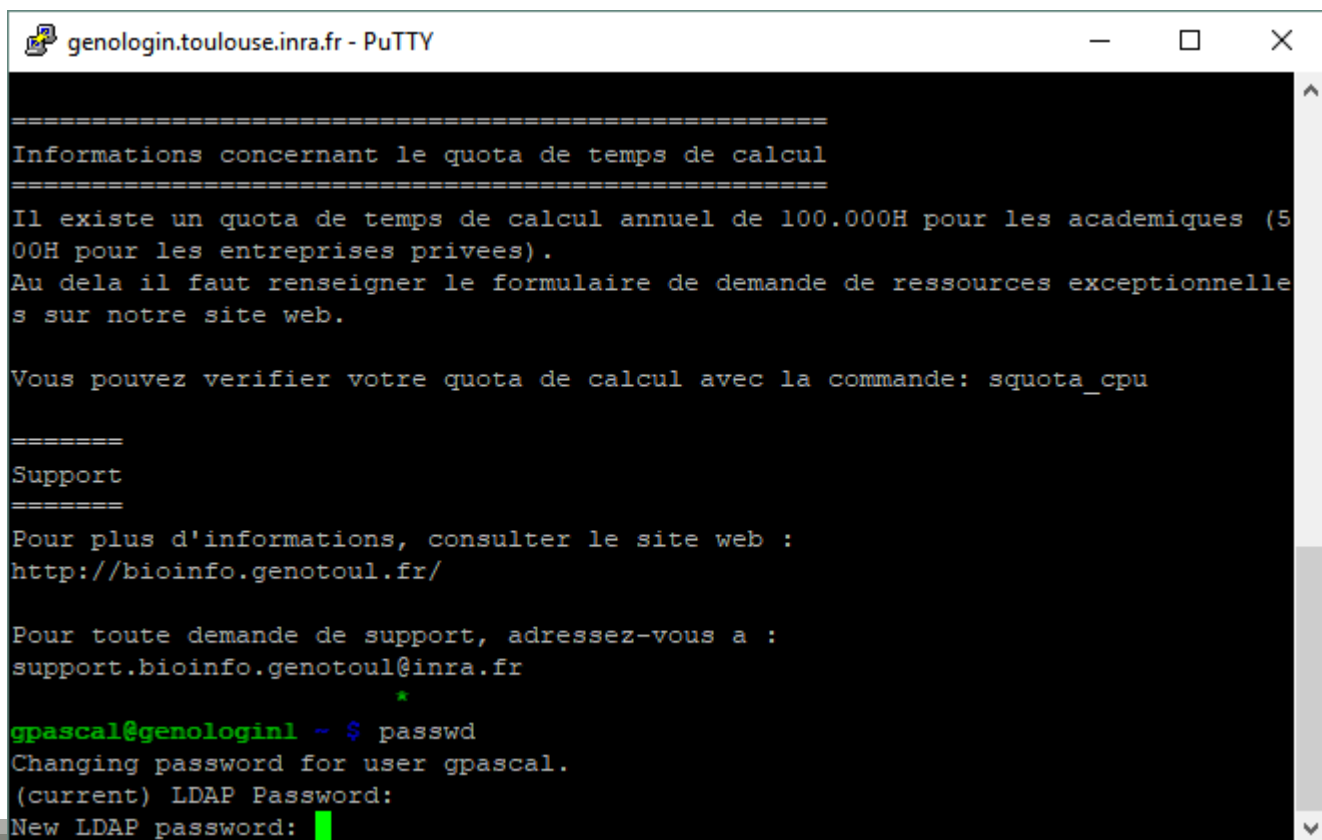
=====
Support
=====

Pour plus d'informations, consulter le site web :
http://bioinfo.genotoul.fr/

Pour toute demande de support, adressez-vous a :
support.bioinfo.genotoul@inra.fr
*
gpascal@genologin1 ~ $ passwd
Changing password for user gpascal.
(current) LDAP Password: █
```

# Change your password

- Enter your new password (with an upper case, a number and a special character in it) and hit « Enter ».



```
genologin.toulouse.inra.fr - PuTTY
=====
Informations concernant le quota de temps de calcul
=====
Il existe un quota de temps de calcul annuel de 100.000H pour les academiques (5
00H pour les entreprises privees).
Au dela il faut renseigner le formulaire de demande de ressources exceptionnelle
s sur notre site web.

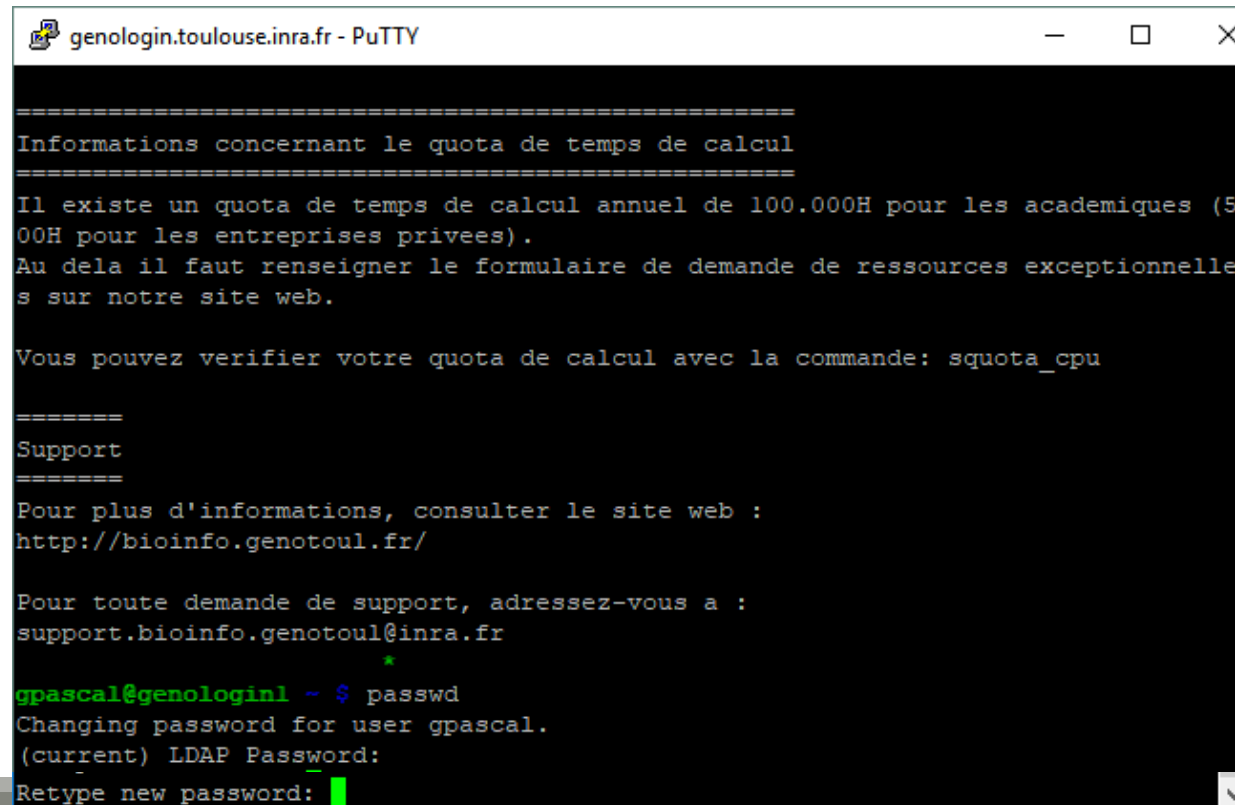
Vous pouvez verifier votre quota de calcul avec la commande: squota_cpu

=====
Support
=====
Pour plus d'informations, consulter le site web :
http://bioinfo.genotoul.fr/

Pour toute demande de support, adressez-vous a :
support.bioinfo.genotoul@inra.fr
*
gpascal@genologin1 ~ $ passwd
Changing password for user gpascal.
(current) LDAP Password:
New LDAP password: █
```

# Change your password

- Enter a second time your new password and hit « Enter ». You have changed your Genotoul password, you can close PuTTY.



```
genologin.toulouse.inra.fr - PuTTY
=====
Informations concernant le quota de temps de calcul
=====
Il existe un quota de temps de calcul annuel de 100.000H pour les academiques (5
00H pour les entreprises privees).
Au dela il faut renseigner le formulaire de demande de ressources exceptionnelle
s sur notre site web.

Vous pouvez verifier votre quota de calcul avec la commande: squota_cpu

=====
Support
=====
Pour plus d'informations, consulter le site web :
http://bioinfo.genotoul.fr/

Pour toute demande de support, adressez-vous a :
support.bioinfo.genotoul@inra.fr

gpascal@genologin1 ~ $ passwd
Changing password for user gpascal.
(current) LDAP Password:
Retype new password: █
```

Galaxy
Analyze Data Workflow Shared Data Visualization Help User
Using 0%

Tools

search tools

MANAGE YOUR DATA FILES

[Get Data](#)

[Download Data](#)

[Jobs statistics](#)

FILES MANIPULATION

[Text Manipulation](#)

[Filter and Sort](#)

[Join, Subtract and Group](#)

[GFF](#)

[BED Tools](#)

[Convert Formats](#)

[Fetch Sequences](#)

Statistics

[Graphical Analysis](#)

SEQUENCES MANIPULATION

[FASTA manipulation](#)

[FastQC: fastq/sam/bam](#)

[Illumina fastq](#)

[Generic FASTQ manipulation](#)

[FASTX-Toolkit for FASTQ data](#)

SAM/BAM MANIPULATION : PICARD (BETA)

[Conversion](#)

[QC/Metrics for sam/bam](#)

[BAM/SAM Cleaning](#)

[SAM/BAM manipulation: SAMtools](#)

[Sequences Queries](#)

[VCF Tools](#)

SGS MAPPING

[BWA - Bowtie](#)

[Indel Analysis](#)

[Variant calling](#)

[SNP annotation](#)

[RNAseq Alignment](#)

[RNAseq Raw Expression](#)

[RNAseq Cufflinks](#)



## How to cite Galaxy workbench ?

Depending on the help provided you can cite us in acknowledgements, references or both.

Examples :

Research teams can thank the Toulouse Midi-Pyrenees bioinformatics platform and Sigeneae group, using in their publications the following sentence : "We are grateful to the genotoul bioinformatics platform Toulouse Midi-Pyrenees and Sigeneae group for providing help and/or computing and/or storage ressources thanks to Galaxy Instance <http://sigeneae-workbench.toulouse.inra.fr>".

In cases of collaboration, you can directly quote the person who participated to the project : Name, Sigeneae group, GenPhySE, INRA Auzeville CS 52627 31326 Castanet Tolosan cedex.

References

X. SIGENAE [<http://www.sigeneae.org/>]

# RESULTS VISUALISATION —AND— TOOL PARAMETER WINDOW



**Ready to use Workflows**

- 1/ FROGS - Find, Rapidly, Otus with Galaxy Solution
- 2/ Tax4Fun workflow
- 3/ Function Table for Tax4Fun matrix workflow



**Galaxy News**

- 1/ How to use [datasets collections](#) ?
- 2/ How to [upload several files in Galaxy](#) ?
- 3/ New tools : Tax4Fun ([manual](#)), VIP ([manual](#)), Salmon, multiQC.
- 4/ Tools availables for [virologie analyses](#)
- 5/ Virology tools
- 6/ How to use [Function Table tool](#) ?



**E-learning**

<https://inra.classilio.com/Login>

2 e-learning availables : Galaxy initiation and Quality sequences analysis

History

search datasets

Data Collection

3 shown

33 b

3: seq1.fasta

2: seq2.fasta

1: seq3.fasta

## DATASETS HISTORY

# Vocabulary of Galaxy

---

- **Tools:**

- A tool has a function which is explained when you click on it.
- Each Galaxy platform has its own tools.

- **Dataset:**

- A dataset is a file, uploaded to Galaxy by you or produced by a tool.
- Be careful: a dataset has a datatype.

- **History:**

- A tool generates datasets and these datasets are stored in the current history.
- Everything is permanently saved.
- If you log off your computer or browser, it's ok, everything will keep running and be saved!

# Your Turn!

---

DISCOVER GALAXY

# Exercise

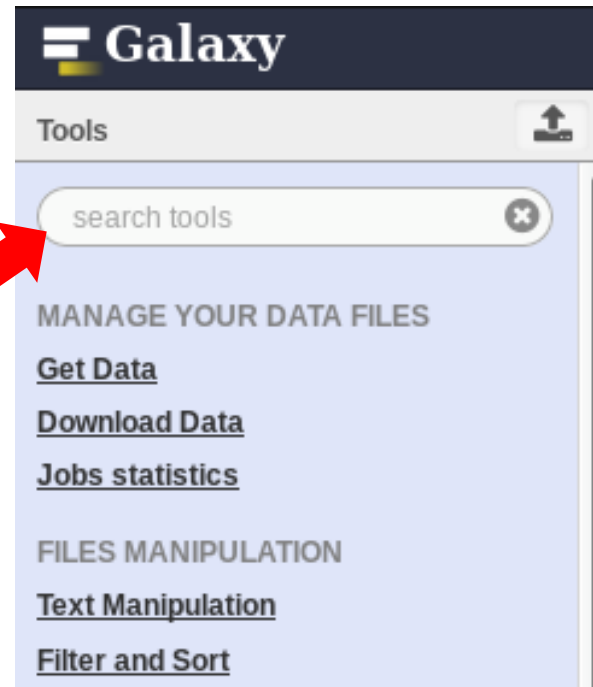
---

1. Visit the Galaxy Platform.
2. Look at the tool list.
3. Display only FROGS tools.
4. Display all tools concerning fastq files.

# Exercise

---

Search a tool by name.





# Manipulate Histories

---

# Your Turn!

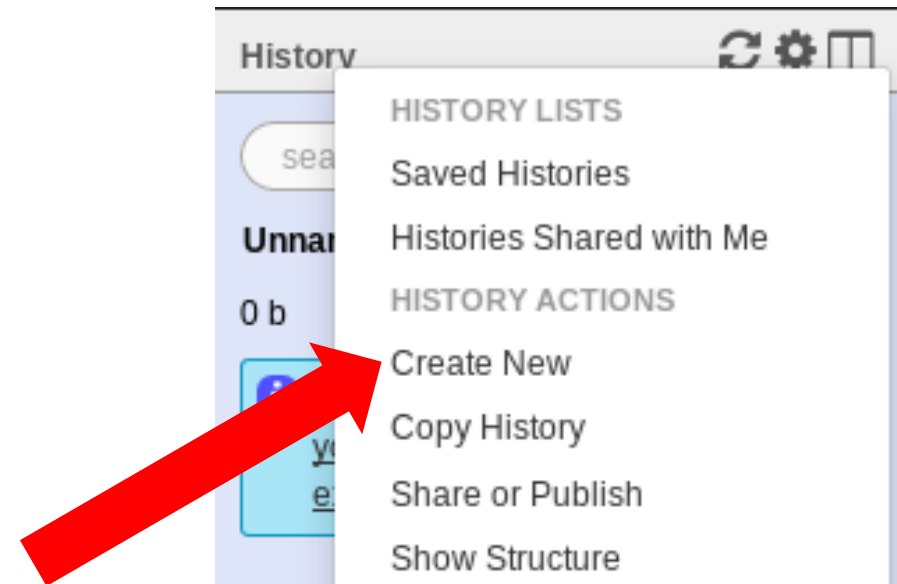
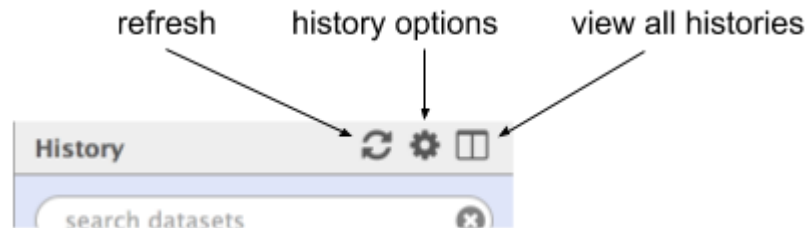
---

CREATE THE 4 HISTORIES NEEDED FOR THE FROGS FORMATION

# To create a new history:

---

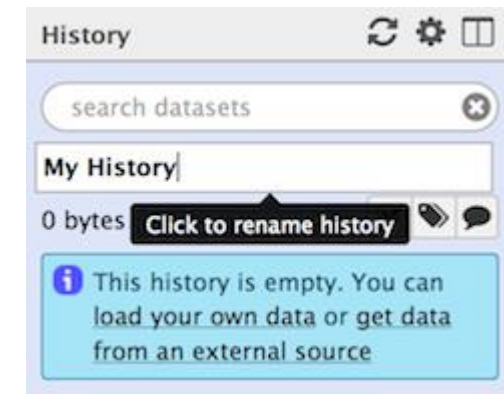
- Click on the wheel.
- Click on « create new ».



# To rename a history:

---

- Click on the history name (at the top).
- Enter « **multiplex** ».
- Hit « **Enter** » to validate.



**Don't use special  
characters or  
accents!**

# Exercise

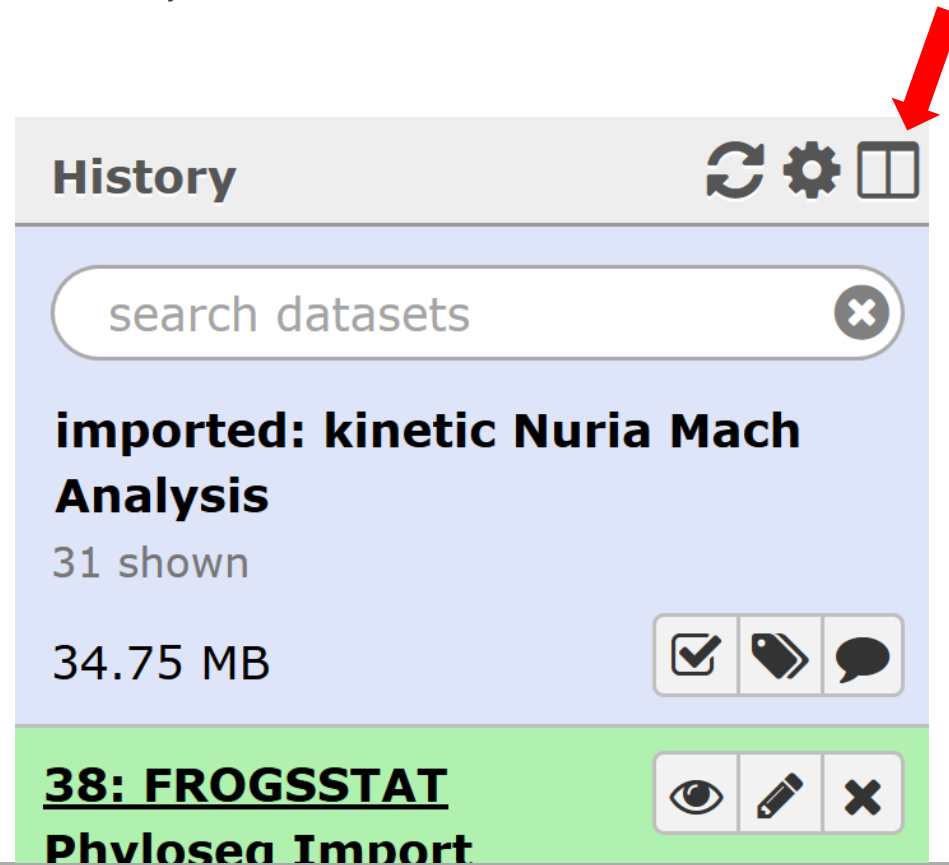
---

- Create histories named:
  - 454 or ITS
  - merged
  - temp
- Switch to the history named « multiplex » as current history.
- Go back to the main interface.

# How to list all histories?

---

- To view all histories, click on this icon.



# Explore the « View all histories » section

**Galaxy** Analyze Data Workflow Shared Data Visualization Help User

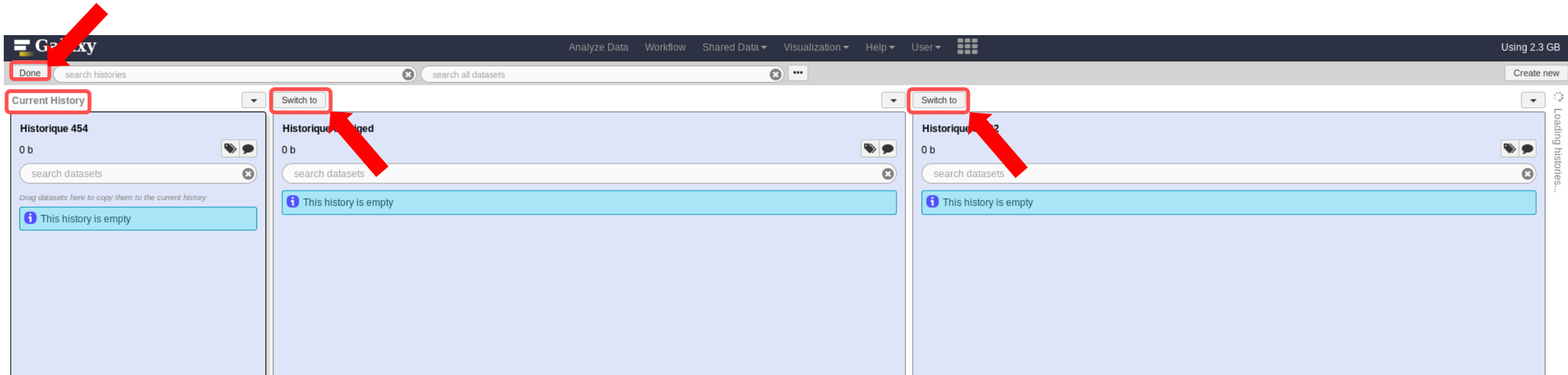
Done search histories search all datasets Create new

**Current History**

- imported: kinetic Nuria Mach Analysis**  
31 shown  
34.75 MB  
search datasets
- Test Ta Thi's tools**  
73 shown  
233.76 MB  
search datasets  
76: FROGS Pre-process: report.html
- Copy of 'full p... maria.bernard...'**  
14 shown  
11.32 MB  
search datasets  
14: FROGS Phv...

38: FROGSSTAT Phylogen...

# Switch current history



- Switch to the history named « multiplex » as current history.
- Click on “Done” to go back to the main interface.



# Data import

---

# How to import your data to Galaxy

---

- 5 ways to upload your data to Galaxy:

- From your computer



- By URL



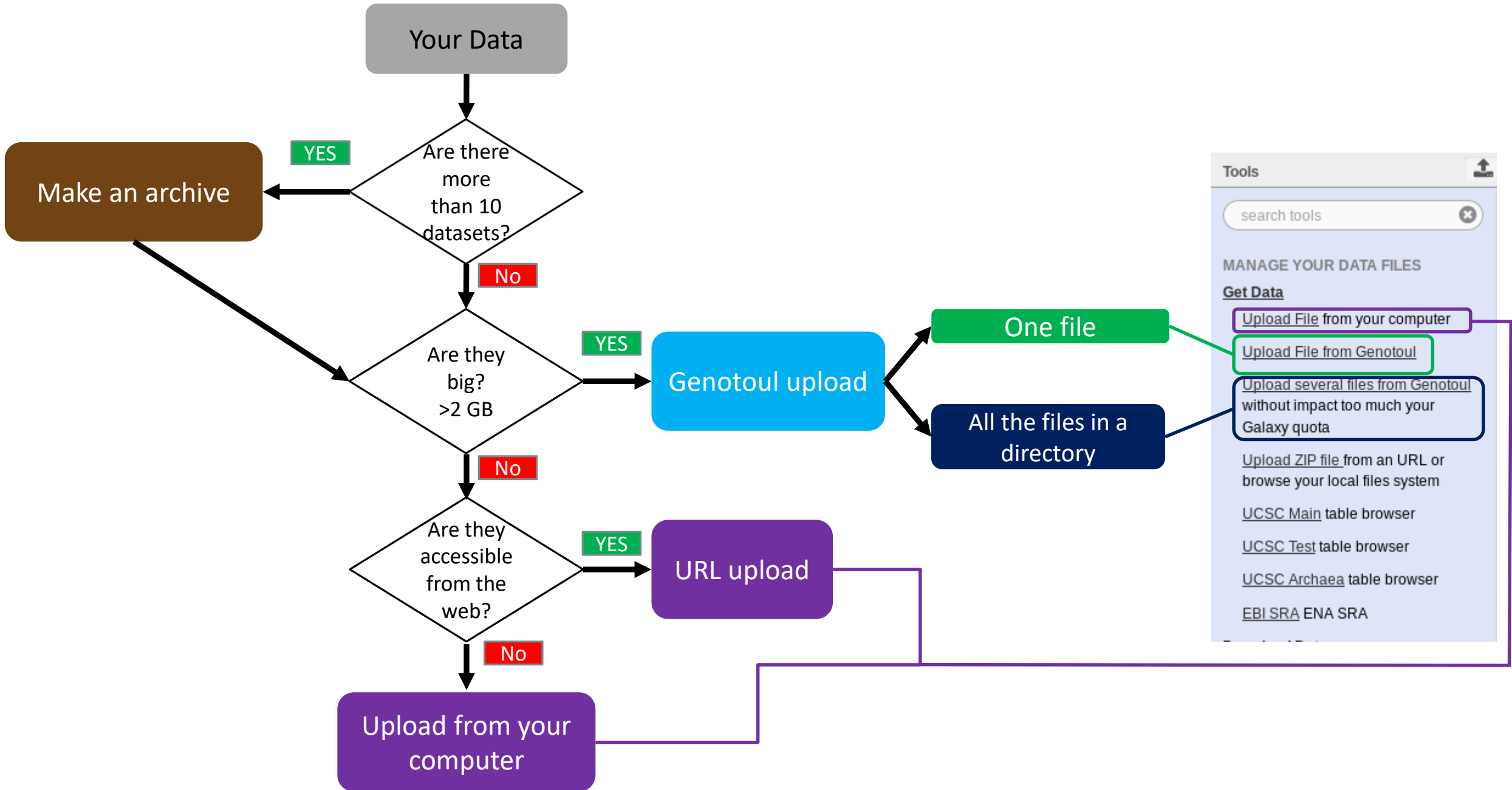
- From Genotoul Bioinfo clusters



- Shared by other users of Galaxy



# How to choose your upload method?



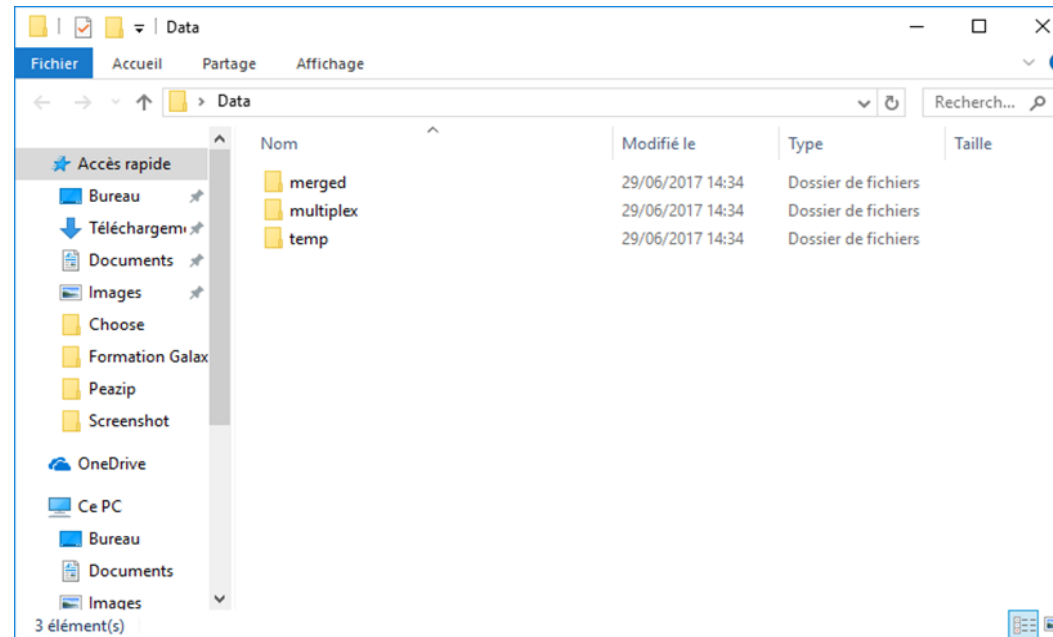
# Your Turn!

---

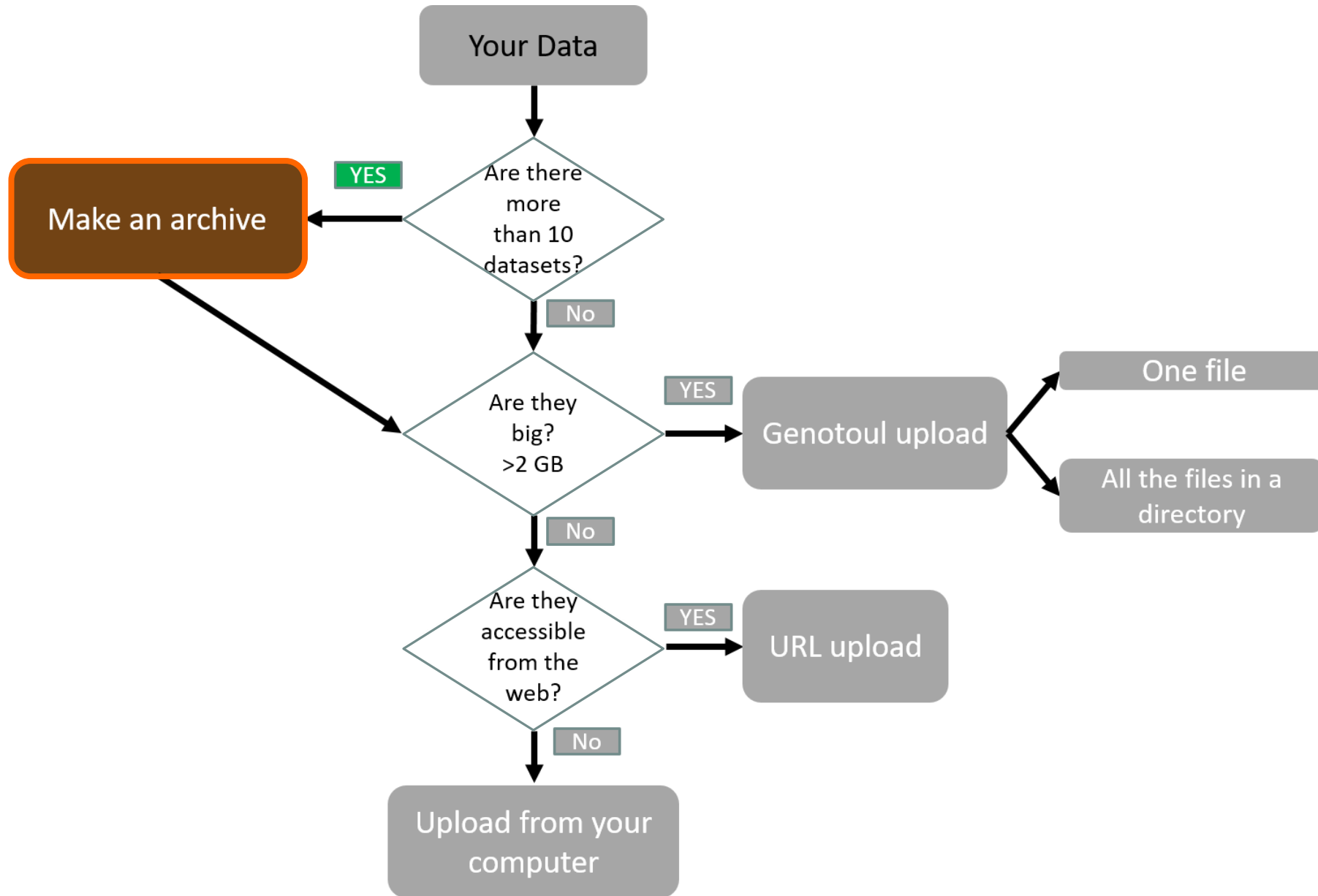
PREPARE FILES

# Formation data

- All the formation data are in the Data folder on your desktop.



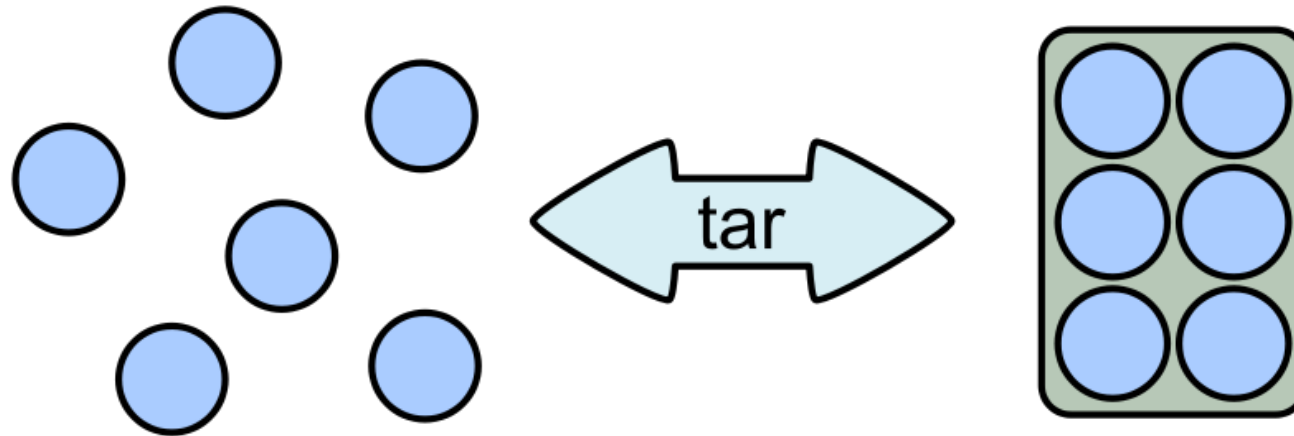
- If you don't have this directory on your desktop, ask us.
- You can find all the formation data online on:  
[http://genoweb.toulouse.inra.fr/~formation/15\\_FROGS/FROGS\\_ini/DATA/Dataset/](http://genoweb.toulouse.inra.fr/~formation/15_FROGS/FROGS_ini/DATA/Dataset/)



# Create a Tar file

---

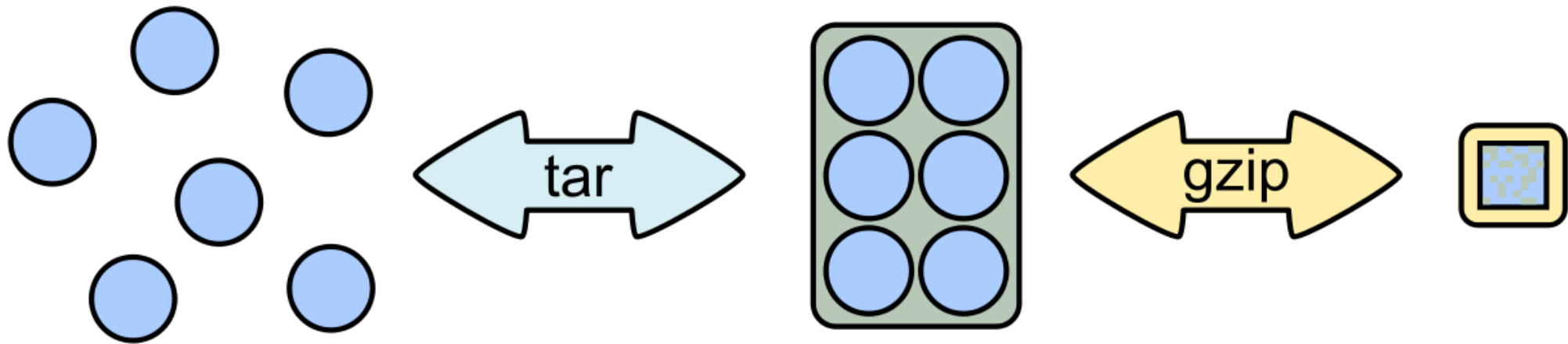
- Uploading multiple local files is time consuming.
- Solution: put all your files in an archive!
- What is a tar file?



# Create a Tar.gz

---

- Moreover, we can compress the archive to free up space.





# Your Turn!

---

CREATE AN ARCHIVE WITH THE FILES IN MERGED

# 7-Zip

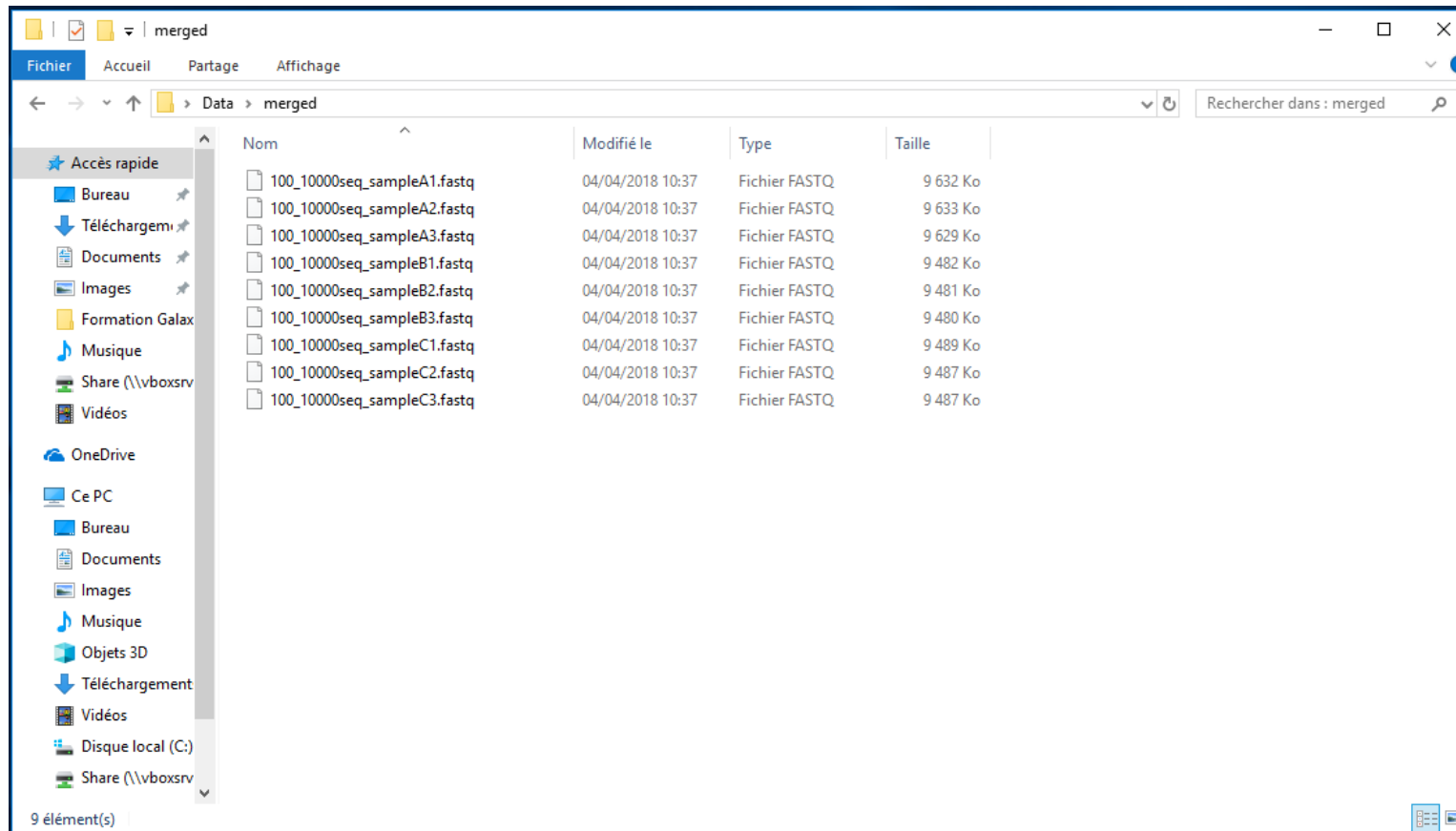
---

- 7-Zip is a software called a file archiver.
- Can archive and compress files.
- Open source
- At the lab, you could download it at: <http://www.7-zip.org/>



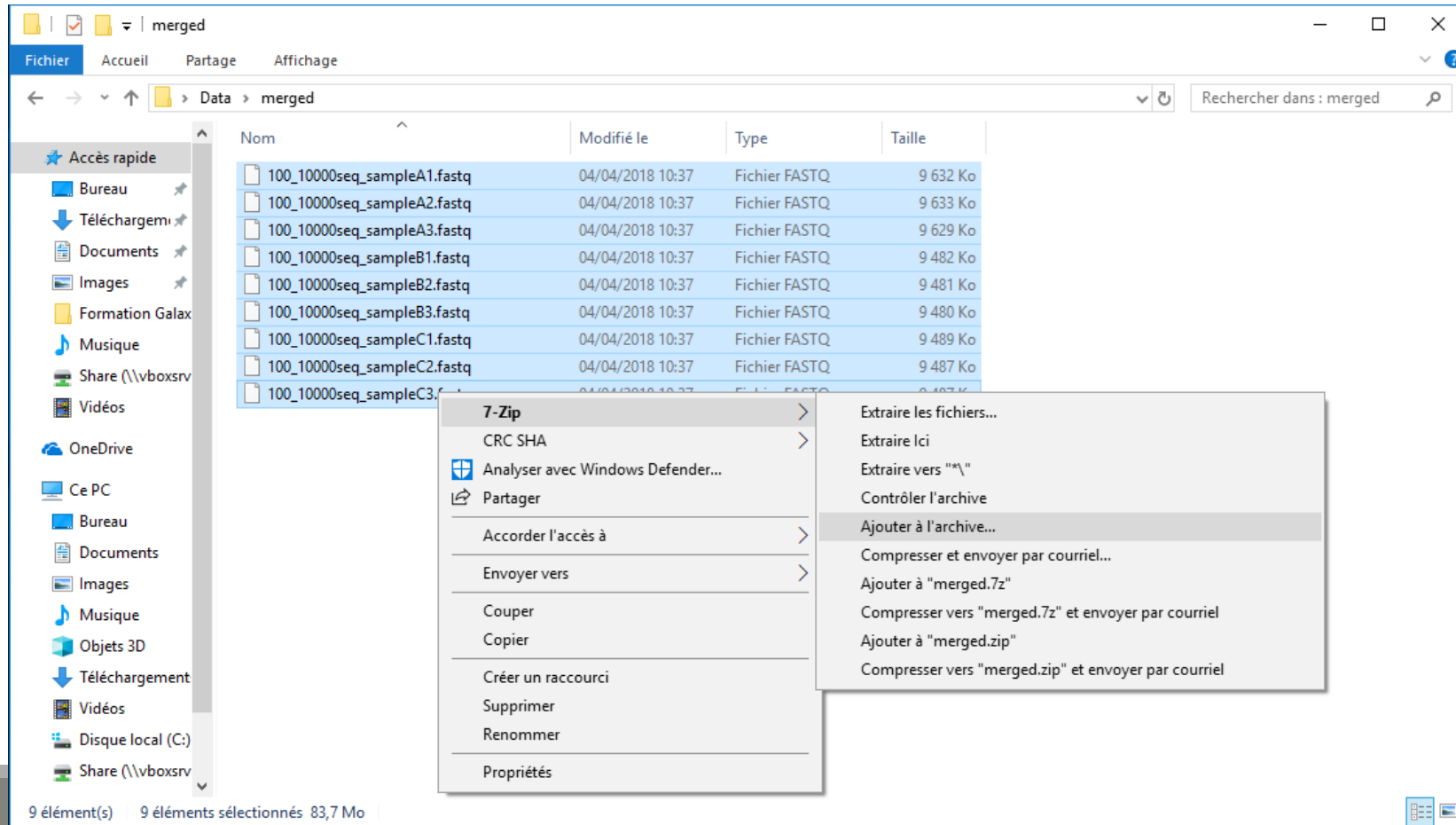
# Create a Tar file with 7-Zip

- Browse into the Data directory and next into the merged directory.



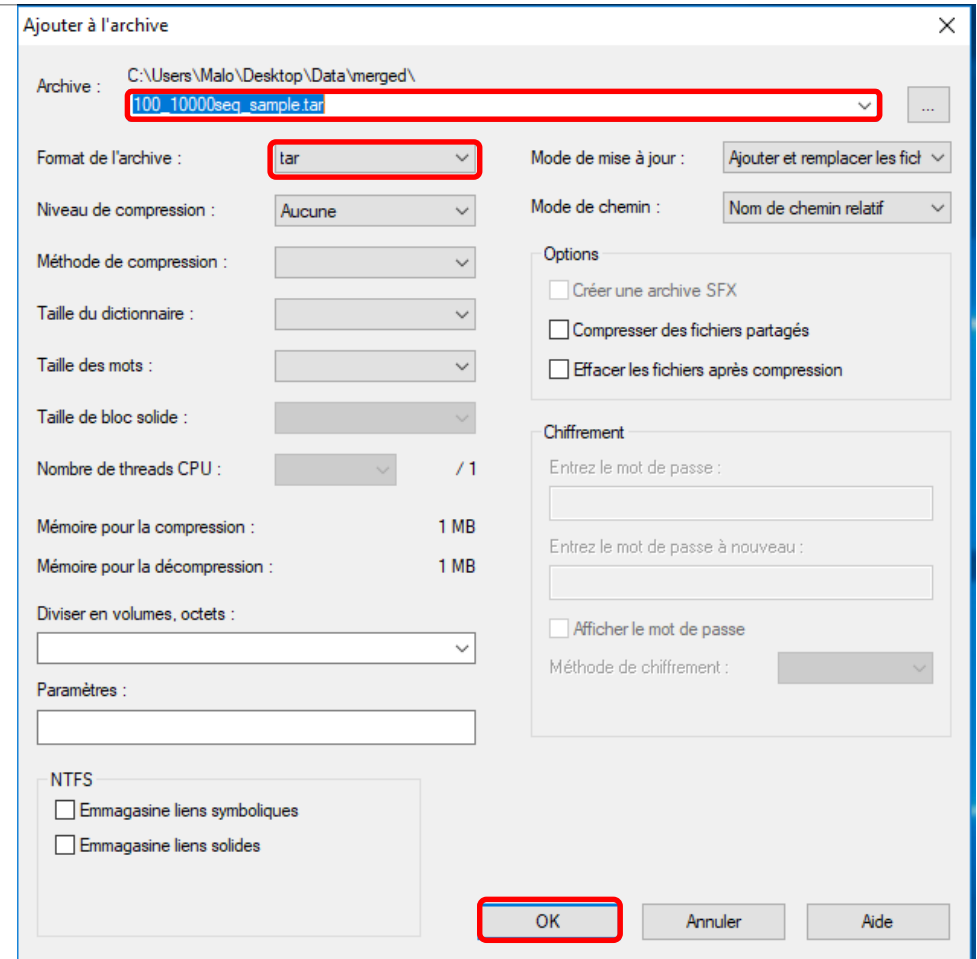
# Create a Tar file with 7-Zip

- Select all files and right click on it > 7-Zip > Add to archive . . .



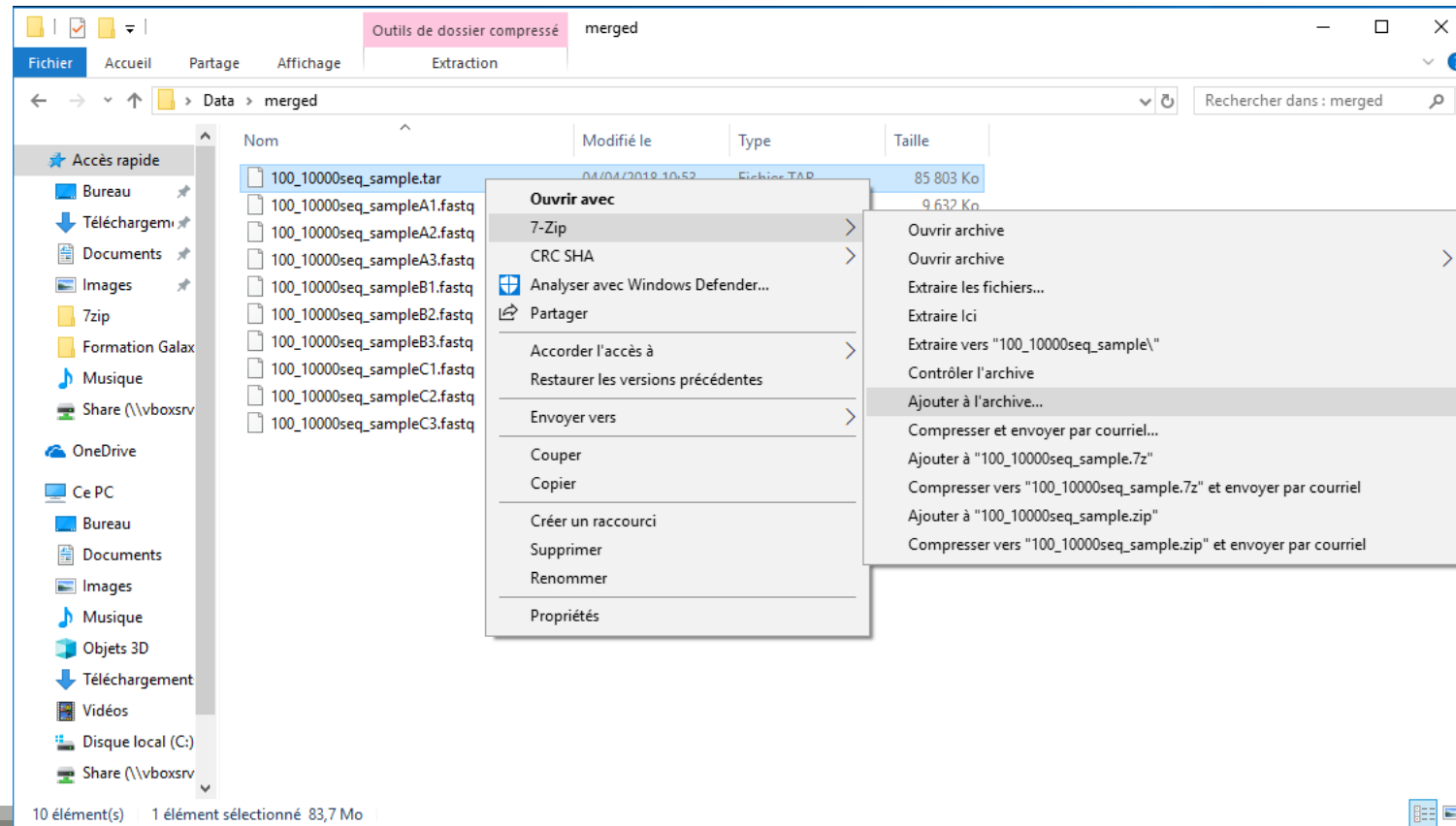
# Create a Tar file with 7-Zip

- Change the name of the archive to:  
« 100\_10000seq\_sample.tar »
- Choose « tar » as archive format .
- Click on OK .



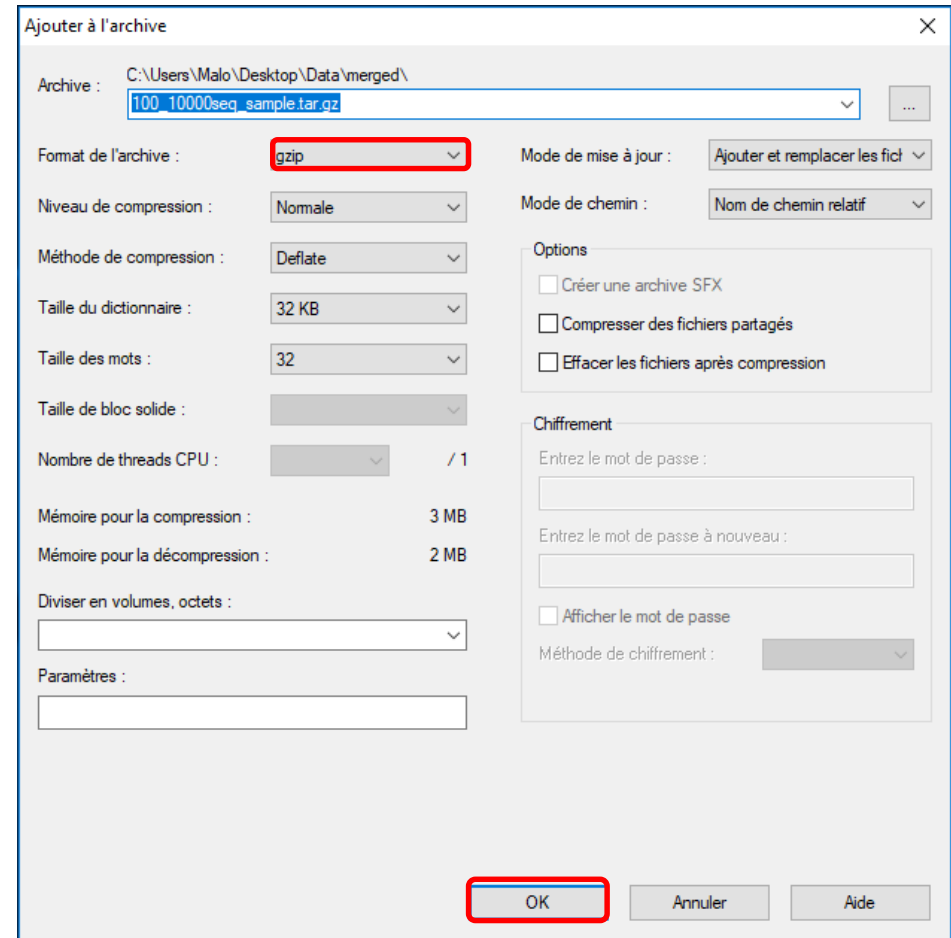
# Create a Tar file with 7-Zip

- Select the newly created archive and right click on it > 7-Zip > Add to archive . . .

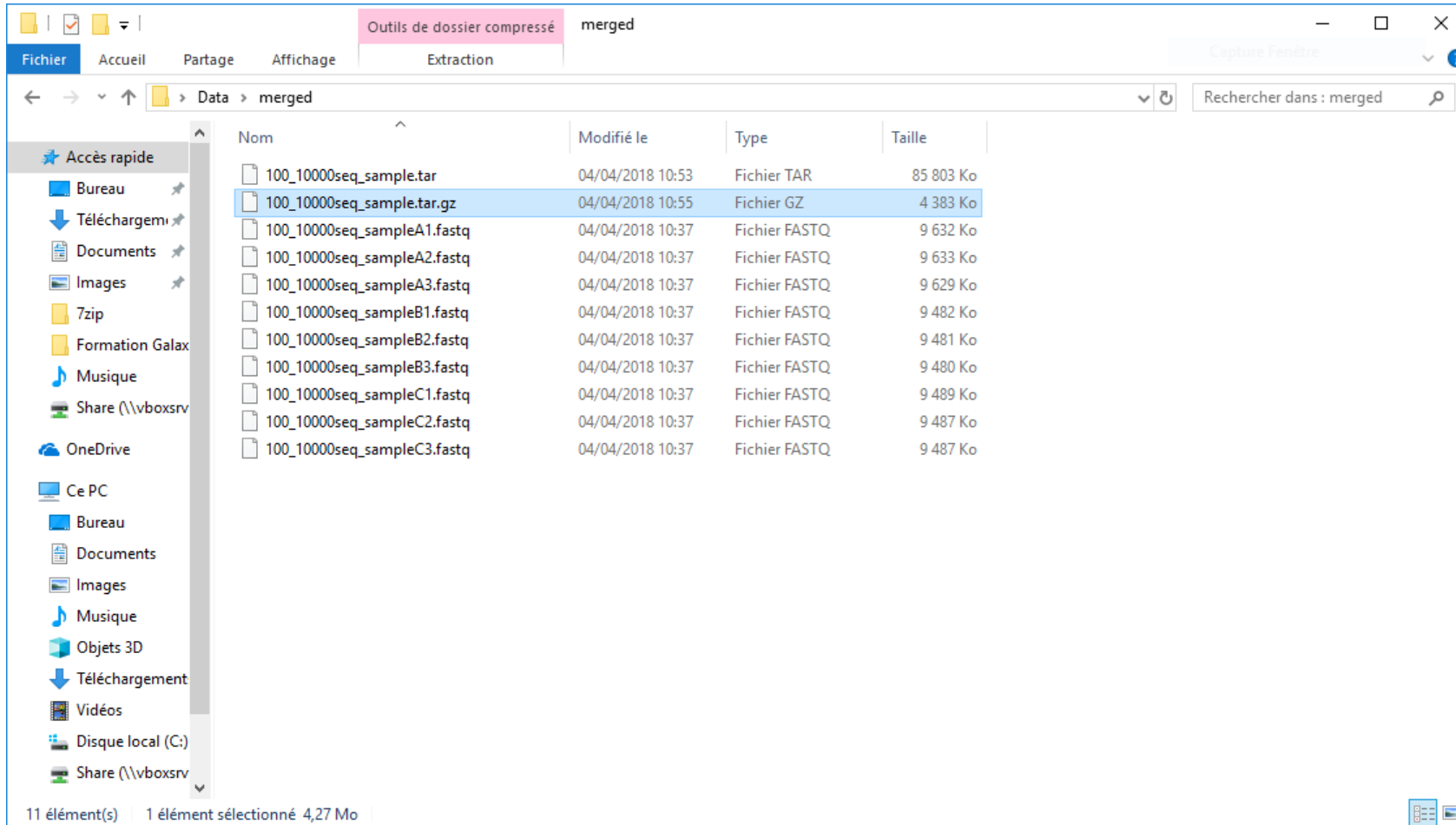


# Create a Tar file with 7-Zip

- Choose « gzip » as archive format .
- Click on OK .

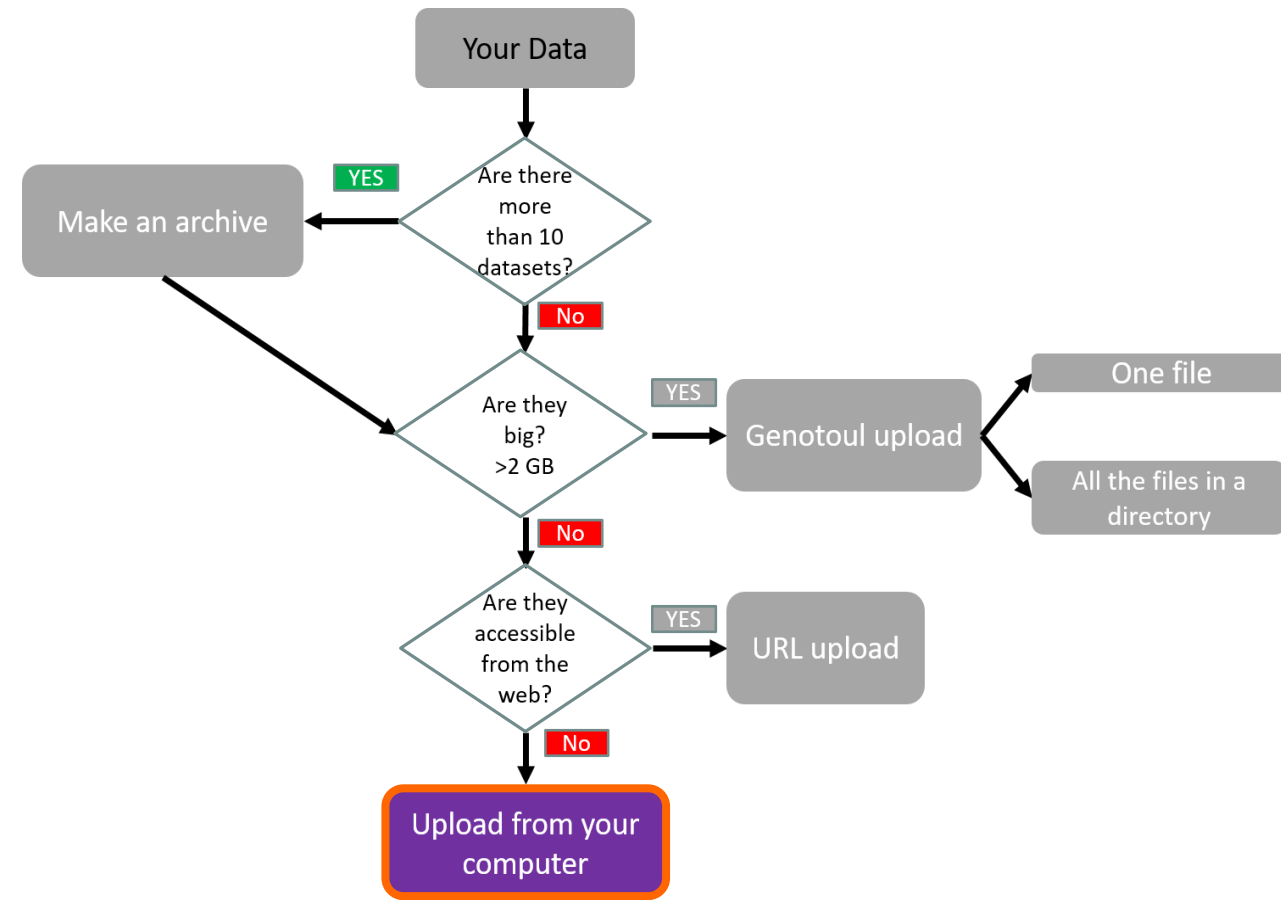


# Create a Tar file with 7-Zip





# Your Turn!

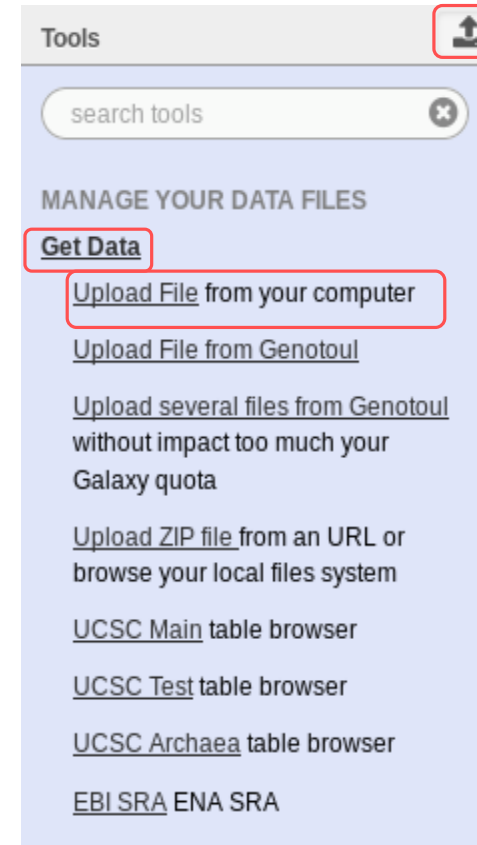


UPLOAD FILES FROM YOUR COMPUTER

# Upload local files

- In Galaxy, your current history must be multiplex.
- Click on the « Get Data » tool.
- Upload file from your computer.

⚠ For files smaller than 2 GB



# Upload local files

Download from web or upload from disk

Regular

Composite

Drop files here

Click here to choose a file on your hard drive.

Upload one of the two files in the multiplex directory.

Type (set all):

Auto-detect



Genome (set all):

unspecified (?)

Choose local file

Paste/Fetch data

Pause

Reset

Start

Close

# Upload local files

The datatype of a file indicates the content of it.

.fasta = fasta

.fastq = fastq

.tar = tar







Select the type corresponding to your file.

Do not trust blindly the auto-detect!


## Download from web or upload from disk

[Regular](#) [Composite](#)

You added 2 file(s) to the queue. Add more files or click 'Start' to proceed.

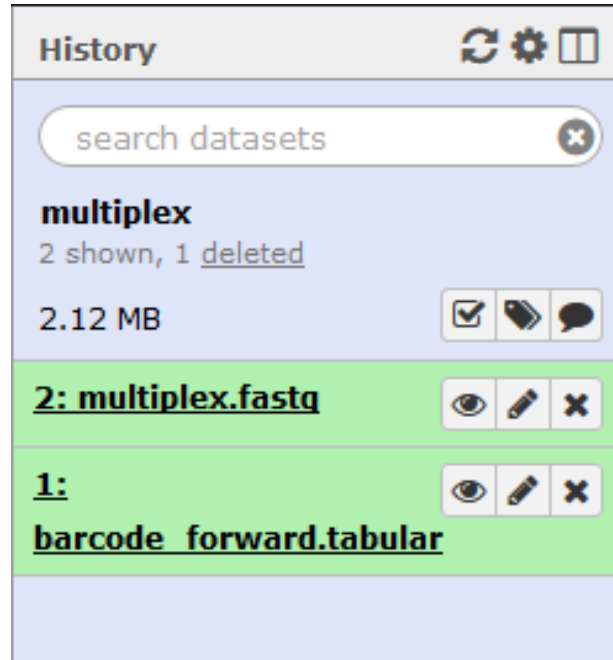
Name	Size	Type	Genome	Settings	Status
 barcode_forward.tabular	260 b	Auto-det...	unspecified (?)		0% 
 multiplex.fastq	2.1 MB	Auto-det...	unspecified (?)		0% 

1. Select the type of file (Do not leave on Auto-Detect!).
2. Select your other files the same way.
3. Begin upload.

Type (set all):   Genome (set all):

# Upload local files

---

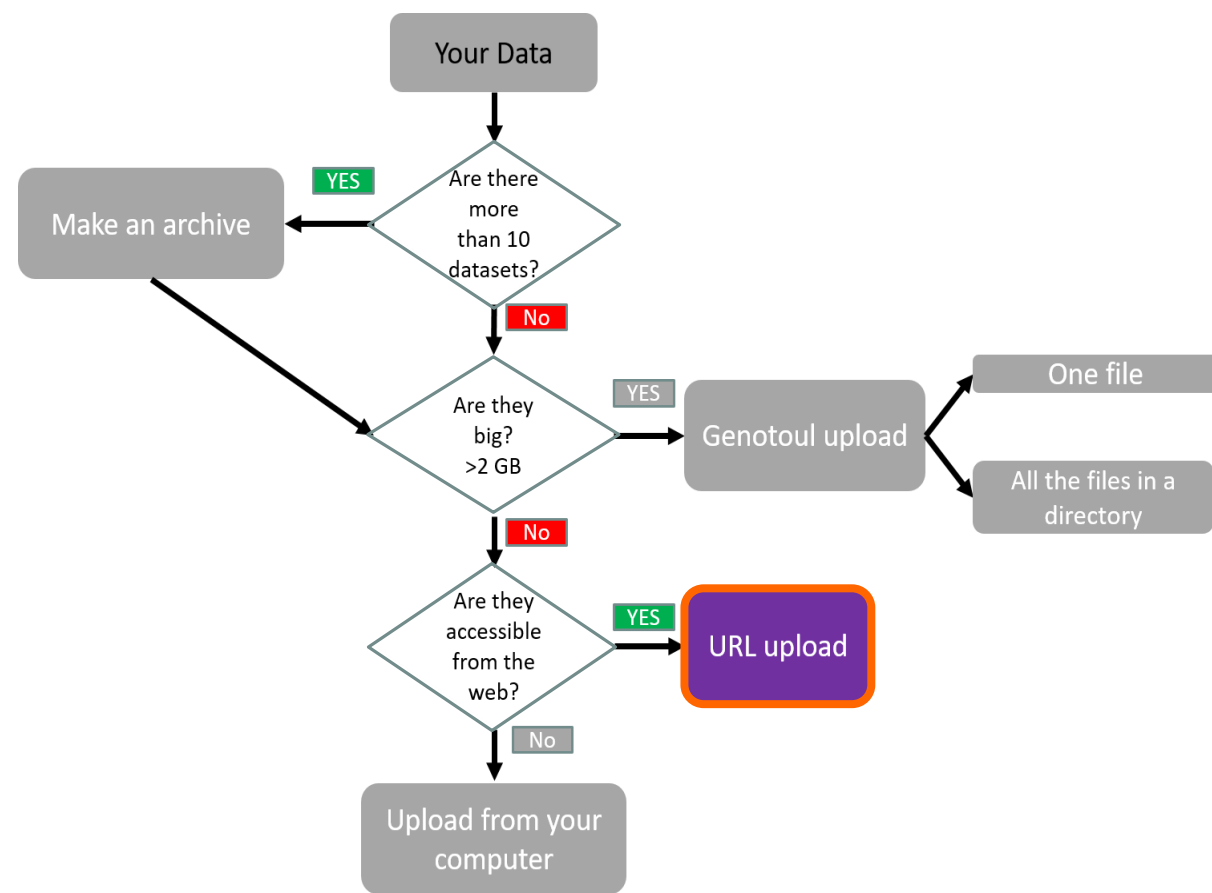


The screenshot shows a 'History' panel with a search bar and a list of datasets. The search bar contains the text 'search datasets'. Below it, the word 'multiplex' is displayed in bold, followed by the text '2 shown, 1 deleted'. The size '2.12 MB' is shown to the left of three icons: a checkmark, a trash can, and a speech bubble. Two dataset entries are listed below, each with its own set of icons (eye, pencil, and X):

- 2: multiplex.fastq**
- 1: barcode forward.tabular**

# Your Turn!

UPLOAD FILE FROM AN URL



# Upload file from URL

1. Switch to 454 or ITS history as current history.
  2. Go to **Get Data > Upload File from your computer**
  3. Click on **Paste/Fetch Data**
  4. Copy the address of the file:  
[http://genoweb.toulouse.inra.fr/~formation/15\\_FROGS/FROGS\\_ini/DATA/454.fastq](http://genoweb.toulouse.inra.fr/~formation/15_FROGS/FROGS_ini/DATA/454.fastq)  
Or  
[http://genoweb.toulouse.inra.fr/~formation/15\\_FROGS/FROGS\\_ini/DATA/ITS.tar.gz](http://genoweb.toulouse.inra.fr/~formation/15_FROGS/FROGS_ini/DATA/ITS.tar.gz)
1. Change the type!
  2. Click on Start.
  3. You can put one address per line for multiple uploads.

## Download from web or upload from disk

Regular Composite

You added 1 file(s) to the queue. Add more files or click 'Start' to proceed.

Name	Size	Type	Genome	Settings	Status
New File	76 b	Auto-detect	unspecified (?)		

You can tell Galaxy to download data from web by entering URL in this box (one per line). You can also directly paste the contents of a file.

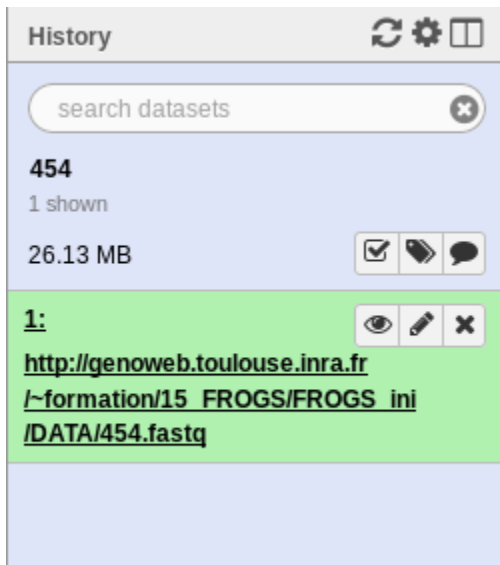
[http://genoweb.toulouse.inra.fr/~formation/15\\_FROGS/FROGS\\_ini/DATA/454.fastq](http://genoweb.toulouse.inra.fr/~formation/15_FROGS/FROGS_ini/DATA/454.fastq)

[http://genoweb.toulouse.inra.fr/~formation/15\\_FROGS/FROGS\\_ini/DATA/ITS.tar.gz](http://genoweb.toulouse.inra.fr/~formation/15_FROGS/FROGS_ini/DATA/ITS.tar.gz)

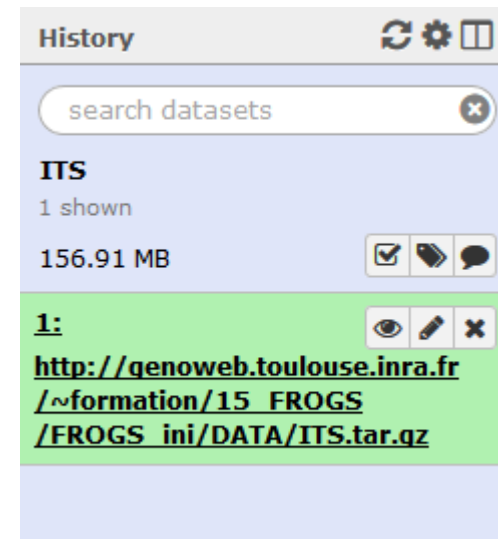
Type (set all): Auto-detect Genome (set all): unspecified (?)

# Upload file from URL

---

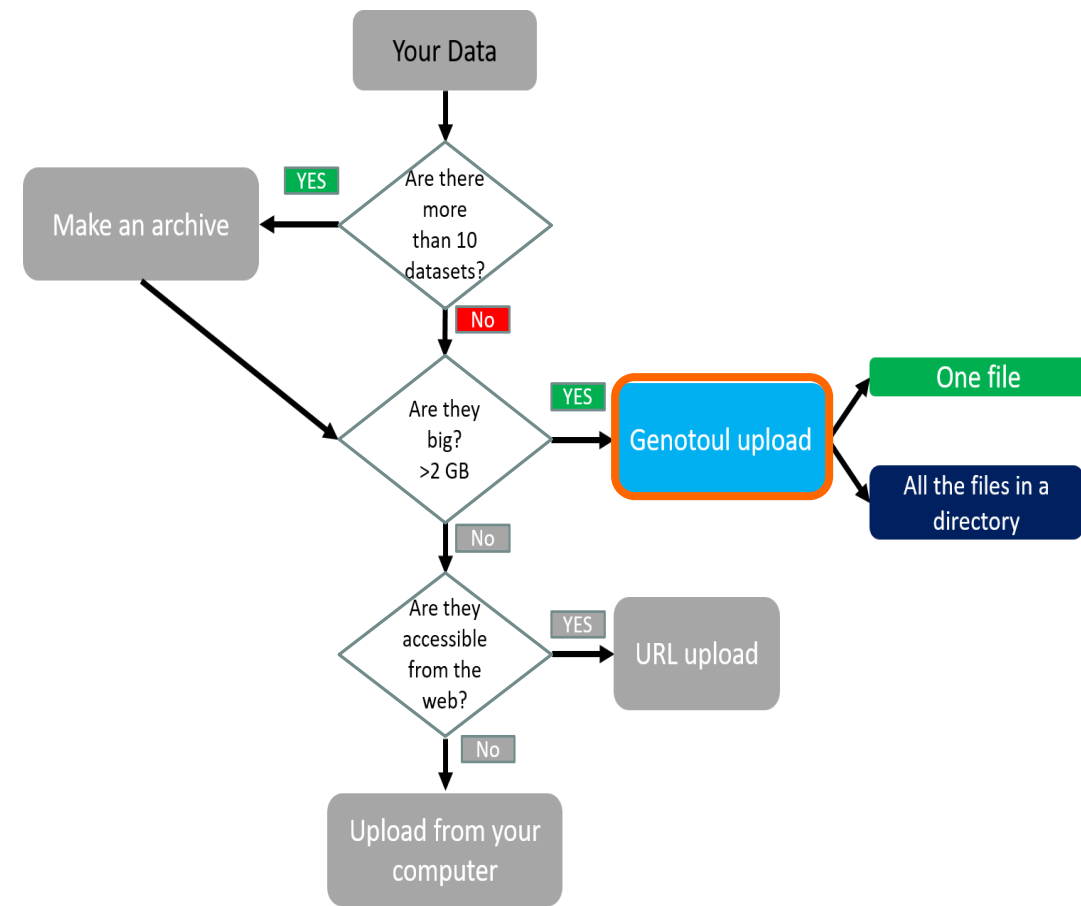


OR



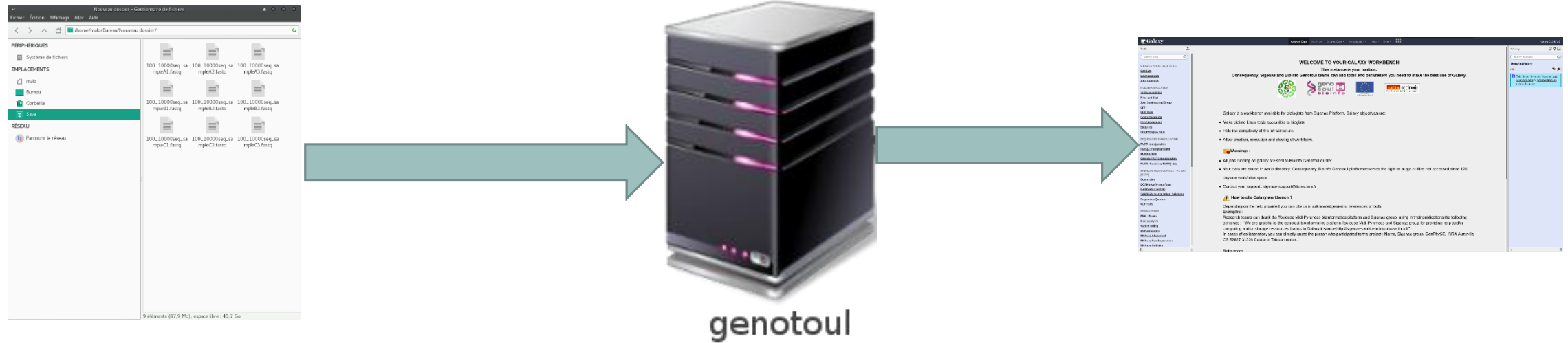


# Your Turn!



UPLOAD FILES TO GENOTOUL AND LINK IT TO GALAXY

# Objectives



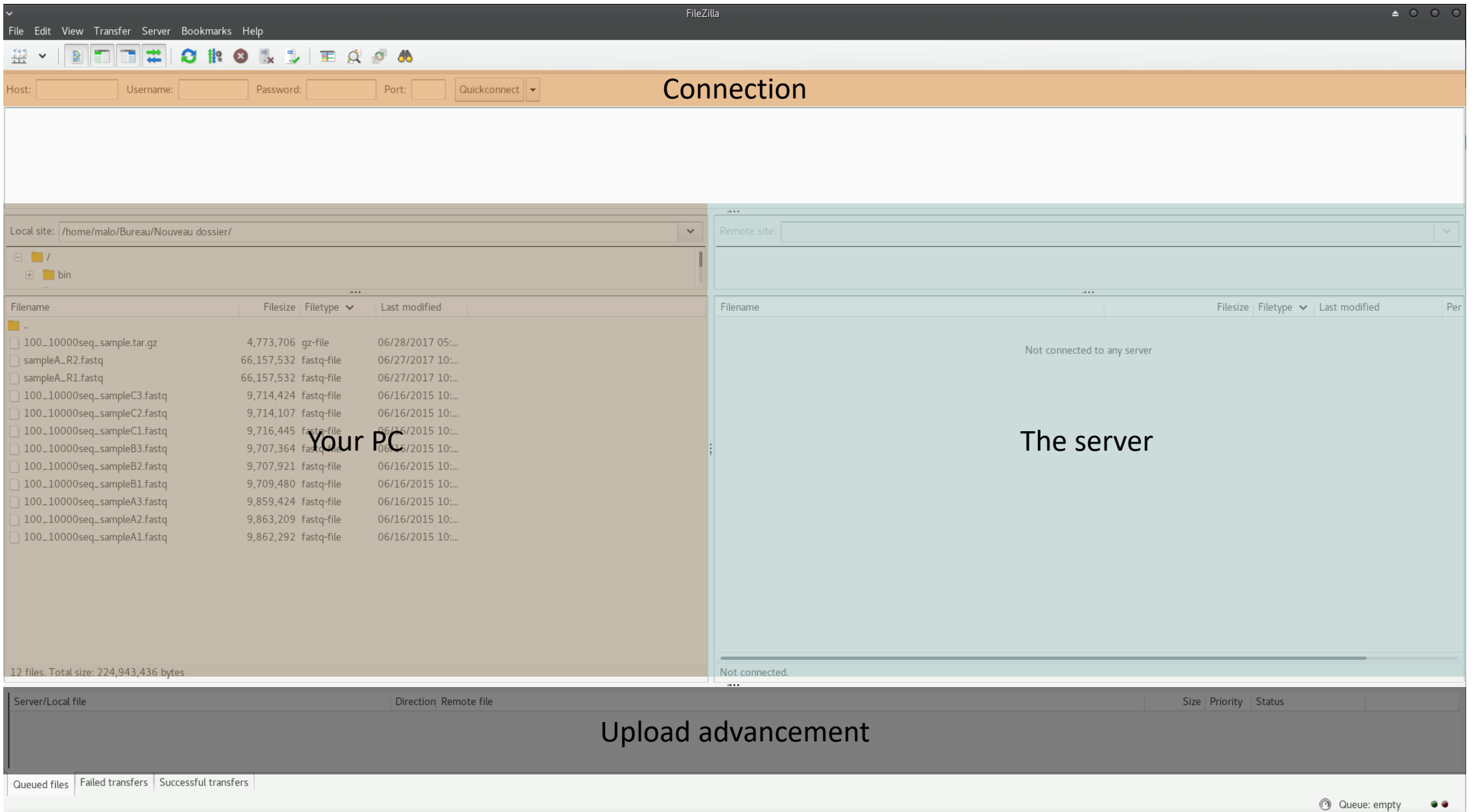
Transfer your files to your Genotoul account and link the file to Galaxy.

# Preparation

---

- Open the data directory on your desktop.
- Launch Filezilla.
  - Filezilla is a FTP client *i.e.* can transfer files to a distant server.
  - You can download it on : <https://filezilla-project.org/>





# Connection

Your PC

The server

# Upload advancement

Host:  Username:  Password:  Port:  Quickconnect

Status: retrieving directory listing of /home/malo/Bureau/work ...  
 Status: Listing directory /work/mleboulch  
 Status: Directory listing of "/work/mleboulch" successful  
 Status: Retrieving directory listing of "/work/mleboulch/Formation" ...  
 Status: Listing directory /work/mleboulch/Formation  
 Status: Directory listing of "/work/mleboulch/Formation" successful  
 Status: Deleting 2 files from "/work/mleboulch/Formation"  
 Status: Disconnected from server

Local site: /home/malo/Bureau/Data/ Remote site: /work/mleboulch/Formation

Local site: /home/malo/Bureau/Data/

- [-] Data
  - merged

Filename	Filesize	Filetype	Last modified
..			
temp		Directory	06/29/2017 05:...
multiplex		Directory	06/29/2017 05:...
merged		Directory	06/29/2017 05:...

3 directories

Remote site: /work/mleboulch/Formation

- [-] mleboulch
  - Formation

Filename	Filesize	Filetype	Last modified	Per
..				
Empty directory listing				

Empty directory.

Browse to the data directory on your desktop.

Server/Local file	Direction	Remote file	Size	Priority	Status

Queued files | Failed transfers | Successful transfers (2)

Queue: empty

Local site: /home/malou/bureau/Data/

Filename	Filesize	Filetype	Last modified
..			
temp		Directory	06/29/2017 05:...
multiplex		Directory	06/29/2017 05:...
merged		Directory	06/29/2017 05:...

3 directories

Remote site: /work/mleboulch/Formation

Filename	Filesize	Filetype	Last modified	Per
..				

Empty directory listing

Empty directory.

Server/Local file | Direction | Remote file | Size | Priority | Status

Queued files | Failed transfers | Successful transfers (2)

Queue: empty

To connect to Genotoul click on this icon.

- Host: genologin.toulouse.inra.fr
- Port: 22
- Protocol: SFTP
- User: your Genotoul login
- Logon Type: Ask for password
- Click on connect.
  
- Password: your password

The screenshot shows the 'Site Manager' dialog box with the 'General' tab selected. The 'Select Entry' pane on the left shows a tree view with 'My Sites' expanded, containing 'Formation' and 'Genotoul'. The 'Genotoul' entry is selected. The main configuration area contains the following fields:

- Protocol: SFTP - SSH File Transfer Protocol
- Host: genologin.toulouse.inra.fr
- Port: 22
- Logon Type: Ask for password
- User: mleboulch
- Password: (empty field)
- Background color: None
- Comments: (empty text area)

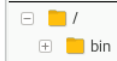
At the bottom of the dialog, there are buttons for 'New Site', 'New Folder', 'New Bookmark', 'Rename', 'Delete', and 'Duplicate'. At the very bottom right, there are buttons for 'Connect', 'OK', and 'Cancel'.

File Edit View Transfer Server Bookmarks Help

Host:  Username:  Password:  Port:  Quickconnect 

Status: Connecting to genotoul.toulouse.inra.fr...  
 Status: Connected to genotoul.toulouse.inra.fr  
 Status: Retrieving directory listing...  
 Status: Listing directory /home/mleboulch  
 Status: Directory listing of "/home/mleboulch" successful

Local site: /home/malo/Bureau/Data/

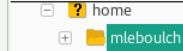


Filename	Filesize	Filetype	Last modified
..			
temp		Directory	06/29/2017 05:...
multiplex		Directory	06/29/2017 05:...
merged		Directory	06/29/2017 05:...

Double click on work directory to access it.



Remote site: /home/mleboulch



Filename	Filesize	Filetype	Last modified
..			
.recently-used.xbel	4,162	xbel-file	03/30/2017 02:26:33 PM
.viminfo	615	File	02/14/2017 09:59:45 AM
.python_history	542	File	06/01/2017 11:08:10 AM
.bashrc	52	File	02/14/2017 09:59:45 AM
.bash_profile	226	File	05/15/2017 03:39:59 PM
.bash_history	17,308	File	06/23/2017 11:15:47 AM
.Xauthority	4,608	File	06/23/2017 11:15:15 AM
work		Directory	10/21/2016 04:37:06 PM
save		Directory	10/21/2016 04:37:06 PM
AIC-prefs		Directory	02/14/2017 10:04:44 AM
.ssh		Directory	01/10/2017 02:28:16 PM
.pki		Directory	03/30/2017 02:56:02 PM
.pathway-tools		Directory	06/01/2017 10:45:23 PM
.gnome2		Directory	02/01/2017 06:14:17 PM
.gconfd		Directory	03/30/2017 02:26:51 PM
.gconf		Directory	03/30/2017 09:08:21 AM
.dbus		Directory	02/01/2017 06:12:40 PM
.config		Directory	04/25/2017 04:13:46 PM

Selected 1 directory.

Server/Local file

Direction Remote file

Size Priority Status

Queued files Failed transfers Successful transfers

Queue: empty



sftp://mleboulch@genotoul.toulouse.inra.fr - FileZilla

File Edit View Transfer Server Bookmarks Help

Host: Username: Password: Port: Quickconnect

Status: Deleting 2 files from /work/mleboulch/Formation  
Status: Disconnected from server  
Status: Retrieving directory listing of "/work/mleboulch" ...  
Status: Listing directory /work/mleboulch  
Status: Directory listing of "/work/mleboulch" successful  
Status: Retrieving directory listing of "/work/mleboulch/Formation" ...  
Status: Listing directory /work/mleboulch/Formation  
Status: Directory listing of "/work/mleboulch/Formation" successful

Local site: /home/malo/Bureau/Data/ Remote site: /work/mleboulch/

Local site: Data merged

Remote site: mleboulch Formation

Filename	Filesize	Filetype	Last modified
..		Directory	06/29/2017 05:...
temp			
multiplex			
merged			

3 directories

Empty directory listing

- Download
- Add files to queue
- View/Edit
- Create directory**
- Create directory and enter it
- Create new file
- Refresh
- Delete
- Rename
- Copy URL(s) to clipboard
- File permissions...

Empty directory.

Server/Local file Direction Remote file Size Priority Status

Queued files Failed transfers Successful transfers (2)

Queue: empty

▪ Create new directory with right click > Create directory.

- Name it « Formation ».
- Go inside this directory next.



sftp://mleboulch@genotoul.toulouse.inra.fr - FileZilla

File Edit View Transfer Server Bookmarks Help

Host: Username: Password: Port: Quickconnect

Status: Listing directory /home/mleboulch  
Status: Directory listing of "/home/mleboulch" successful  
Status: Retrieving directory listing of "/home/mleboulch/work"...  
Status: Listing directory /work/mleboulch  
Status: Directory listing of "/work/mleboulch" successful  
Status: Retrieving directory listing of "/work/mleboulch/Formation"...  
Status: Listing directory /work/mleboulch/Formation  
Status: Directory listing of "/work/mleboulch/Formation" successful

Local site: /home/malo/Bureau/Data/ Remote site: /work/mleboulch/Formation

Local site tree: / bin

Filename	Filesize	Filetype	Last modified
..			
temp		Directory	06/29/2017 05:...
multiplex		Directory	06/29/2017 05:...
merged		Directory	06/29/2017 05:...

Remote site tree: mleboulch Formation

Filename	Filesize	Filetype	Last modified	Per
..				

Empty directory listing

Selected 1 directory. Empty directory.

Server/Local file	Direction	Remote file	Size	Priority	Status
-------------------	-----------	-------------	------	----------	--------

Queued files Failed transfers Successful transfers

Queue: empty

- Select the temp directory and drag and drop it following the arrow.
- Double click on the merge directory.

Host:  Username:  Password:  Port:  Quickconnect 

```
Status: Listing directory /work/mleboulch/Formation/temp
Command: put "/home/malo/Bureau/Data/temp/temp2.fastq" "temp2.fastq"
Command: local:/home/malo/Bureau/Data/temp/temp2.fastq => remote:/work/mleboulch/Formation/temp/temp2.fastq
Status: File transfer successful, transferred 2,226,378 bytes in 1 second
Status: File transfer successful, transferred 2,226,378 bytes in 1 second
Status: Retrieving directory listing of "/work/mleboulch/Formation"...
Status: Listing directory /work/mleboulch/Formation
Status: Directory listing of "/work/mleboulch/Formation" successful
```

Local site: /home/malo/Bureau/Data/merged/

▾ Data  
    ▾ merged

Filename	Filesize	Filetype	Last modified
..			
100_1000seq_sample.tar.gz	4,773,706	gz-file	06/28/2017 05:...
100_1000seq_sampleC3.fastq	9,714,424	fastq-file	06/16/2015 10:...
100_1000seq_sampleC2.fastq	9,714,107	fastq-file	06/16/2015 10:...
100_1000seq_sampleC1.fastq	9,716,445	fastq-file	06/16/2015 10:...
100_1000seq_sampleB3.fastq	9,707,364	fastq-file	06/16/2015 10:...
100_1000seq_sampleB2.fastq	9,707,921	fastq-file	06/16/2015 10:...
100_1000seq_sampleB1.fastq	9,709,480	fastq-file	06/16/2015 10:...
100_1000seq_sampleA3.fastq	9,859,424	fastq-file	06/16/2015 10:...
100_1000seq_sampleA2.fastq	9,863,209	fastq-file	06/16/2015 10:...
100_1000seq_sampleA1.fastq	9,862,292	fastq-file	06/16/2015 10:...

Selected 1 file. Total size: 4,773,706 bytes

Server/Local file

Dire

Queued files | Failed transfers | Successful transfers (2)

Remote site: /work/mleboulch/Formation

▾ mleboulch  
    + ▾ Formation

Filename	Filesize	Filetype	Last modified	Per
..				
temp		Directory	06/29/2017 06:01:32 PM	drw

- Select the 100\_1000seq\_sample.tar.gz file and drag and drop it following the arrow.
- Double click on the merge directory.

Queue: empty


 Host:  Username:  Password:  Port:  Quickconnect 

Status: Directory listing of "/work/.vle" successful  
 Status: Retrieving directory listing of "/work/BPMP" ...  
 Status: Listing directory /work/BPMP  
 Status: Directory listing of "/work/BPMP" successful  
 Status: Retrieving directory listing of "/work/mlebeurier2" ...  
 Command: cd "/work/mlebeurier2"  
 Error: Directory /work/mlebeurier2: permission denied  
 Error: Failed to retrieve directory listing

Local site: /home/malo/Bureau/Data/merged/

Remote site: /work/mleboulch



Filename	Filesize	Filetype	Last modified
100_10000seq_sampleB1.fastq	9,709,480	fastq-file	04/04/2018 10:...
100_10000seq_sampleC2.fastq	9,714,107	fastq-file	04/04/2018 10:...
100_10000seq_sampleC3.fastq	9,714,424	fastq-file	04/04/2018 10:...
100_10000seq_sampleC1.fastq	9,716,445	fastq-file	04/04/2018 10:...
100_10000seq_sampleA3.fastq	9,859,424	fastq-file	04/04/2018 10:...
100_10000seq_sampleA1.fastq	9,862,292	fastq-file	04/04/2018 10:...

Filename	Filesize	Filetype	Last modified	Permis
...		Directory	06/08/2018 11:19:08 AM	drwxr->
...		Directory	03/14/2018 02:21:50 PM	drwxr->
...		Directory	03/13/2018 11:50:00 AM	drwxr->
...		Directory	03/09/2018 04:09:45 PM	drwxr->
...		Directory	03/06/2018 04:25:02 PM	drwxr->
...		Directory	03/06/2018 01:24:27 PM	drwxr->
...		Directory	02/28/2018 03:48:40 PM	drwxr->
...		Directory	02/28/2018 03:47:01 PM	drwxr->
...		Directory	02/28/2018 09:29:46 AM	drwxr->
...		Directory	10/30/2017 05:07:06 PM	drwxr->
...		Directory	06/30/2017 04:26:59 PM	drwxr->

Select your work directory.

Right click on it and click on File Attributes.

Queued files Failed transfers Successful transfers

Queue: empty

Change file attributes

Please select the new attributes for the directory "Formation".

Owner permissions

<input checked="" type="checkbox"/> Read	<input checked="" type="checkbox"/> Write	<input checked="" type="checkbox"/> Execute
--	---	---

Group permissions

<input checked="" type="checkbox"/> Read	<input type="checkbox"/> Write	<input checked="" type="checkbox"/> Execute
--	--------------------------------	---

Public permissions

<input checked="" type="checkbox"/> Read	<input type="checkbox"/> Write	<input checked="" type="checkbox"/> Execute
--	--------------------------------	---

Numeric value:

You can use an x at any position to keep the permission the original files have.

Recurse into subdirectories

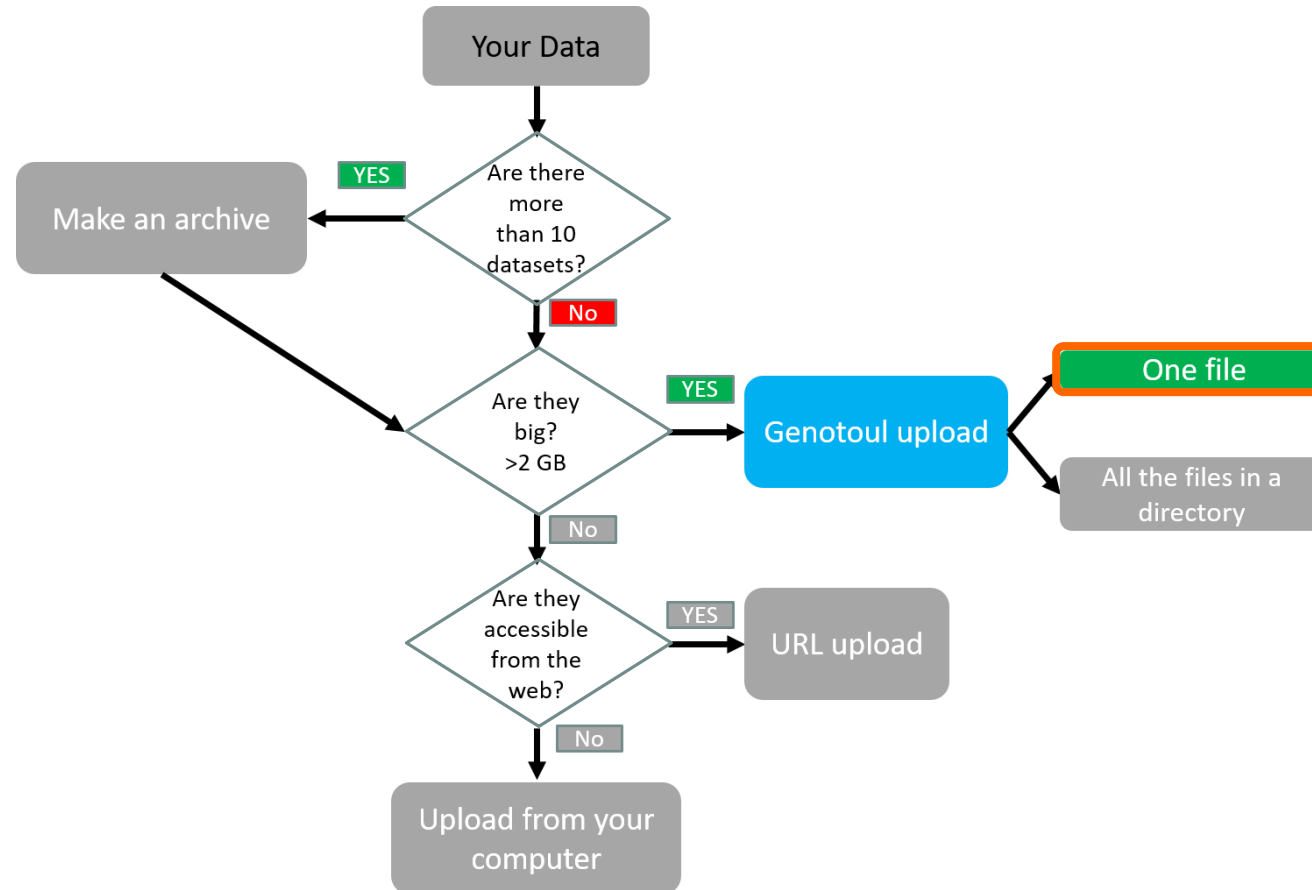
Apply to all files and directories

Apply to files only

Apply to directories only

- Check that all boxes for execute and read are checked.
- Recurse this action to all files and subdirectories.
- That allows Galaxy to access your files on Genotoul.
  - Click on Ok.

# Upload files from Genotoul



File Edit View Transfer Server Bookmarks Help

Host:  Username:  Password:  Port:  Quickconnect

Status: Setting permissions of /work/mleboulch/Formation/temp/temp2.fastq to 755  
Status: /work/mleboulch/Formation/temp/temp2.fastq: 0644 -> 0755  
Status: Retrieving directory listing of "/work/mleboulch/Formation"  
Status: Listing directory /work/mleboulch/Formation  
Status: Directory listing of "/work/mleboulch/Formation" successful  
Status: Retrieving directory listing of "/work/mleboulch/Formation/temp"  
Status: Listing directory /work/mleboulch/Formation/temp  
Status: Directory listing of "/work/mleboulch/Formation/temp" successful

Local site: /home/malo/Bureau/Data/merged/

Remote: /work/mleboulch/Formation

Data  
merged

Filename	Filesize	Filetype	Last modified	Per
..				
100_10000seq_sampleB2.fastq	9,707,921	fastq-file	06/16/2015 10:...	
100_10000seq_sampleB1.fastq	9,709,480	fastq-file	06/16/2015 10:...	
100_10000seq_sampleA3.fastq	9,859,424	fastq-file	06/16/2015 10:...	
100_10000seq_sampleA2.fastq	9,863,209	fastq-file	06/16/2015 10:...	
100_10000seq_sampleA1.fastq	9,862,292	fastq-file	06/16/2015 10:...	

10 files. Total size: 92,628,372 bytes

Formation  
temp

Filename	Filesize	Filetype	Last modified	Per
..				
100_10000seq_sample.tar.gz	4,773,706	gz-file	06/29/2017 06:04:35 PM	-rwx
temp		Directory	06/29/2017 06:01:32 PM	drwx

1 file and 1 directory. Total size: 4,773,706 bytes

Server/Local file | Direction | Remote file | Size | Priority | Status

Queued files | Failed transfers | Successful transfers (3)

Queue: empty

Select this text and copy it by pressing the keyboard keys Ctrl+C.

Tools

search tools

MANAGE YOUR DATA FILES

**Get Data**

- Upload File from your computer
- Upload File from Genotoul
- Upload several files from Genotoul without impact too much your Galaxy quota
- Upload ZIP file from an URL or browse your local files system
- UCSC Main table browser
- UCSC Test table browser
- UCSC Archaea table browser
- EBI SRA ENA SRA

Download Data

Jobs statistics

FILES MANIPULATION

- Text Manipulation
- Filter and Sort
- Join, Subtract and Group
- GFF
- BED Tools
- Convert Formats
- Fetch Sequences
- Statistics
- Graph/Display Data

SEQUENCES MANIPULATION

- FASTA manipulation
- FastQC: fastq/sam/bam
- Illumina fastq
- Generic FASTQ manipulation
- FASTX-Toolkit for FASTQ data

SAM/BAM MANIPULATION : PICARD (BETA)

- Conversion
- QC/Metrics for sam/bam

### Upload File from Genotoul (Galaxy Version 1.0.0)

Options

**Path to file**

Path must be like : /work/USERNAME/somewhere/afile

**File type**

tar.gz

Execute

**What it does**

This tool allows you to use a file stored in your genotoul work directory and optimize Galaxy work space by creating symlinks.

**Path to file**

This must be an absolute path

valid path : /work/LinuxUserName/...  
invalid path : /home/LinuxUserName/...

- Switch to merged history.
- Next go to Get Data > Upload File from Genotoul.

**⚠️ To use this tool and to maintain the confidentiality of yours directories:**

- Create a "galaxy" directory in your work : `mkdir galaxy`
- `chmod a+x /work/LinuxUserName`

Example : `drwxr-x--x 4 smaman sigenae 16384 mar 9 14:15 /work/smaman`

- `chmod a+r /work/LinuxUserName/dataGalaxy.fasta`

**ℹ️ Thanks to the fact that this tool requires you to enter your filepath (without "browse" button), you can manage your data privacy.**

For example, if your data to download in Galaxy are: `/work/LinuxUserName/galaxy/data.fasta`:

- Add "x" rights to "others" on `/work/LinuxUserName/` and on `galaxy/`

It is not useful that "others" have "r" rights of these directories.

- Add "r" rights (only) to "others" on `data.fasta` file.

Thus, Galaxy can access and read `data.fasta` but all other files in those directories will not be accessible or readable.

**⚠️ fastq files have to be uploaded in a correct format (for instance, `fatsqsanger`) in order to be used by Galaxy tools. If this is not the case, your fastq files uploaded will not be listed among available datasets in Galaxy tools.**

---

Version Galaxy Tool : V1.0

Versions of bioinformatics tools used : No bioinformatique tool used.

History

search datasets

merged

0 b

ℹ️ This history is empty. You can [load your own data](#) or [get data from an external source](#)



Tools

search tools

MANAGE YOUR DATA FILES

**Get Data**

- Upload File from your computer
- Upload File from Genotoul
- Upload several files from Genotoul without impact too much your Galaxy quota
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- UCSC Main table browser
- UCSC Test table browser
- UCSC Archaea table browser
- EBI SRA ENA SRA

**Download Data**

**Jobs statistics**

FILES MANIPULATION

- Text Manipulation
- Filter and Sort
- Join, Subtract and Group
- GFF
- BED Tools
- Convert Formats
- Fetch Sequences
- Statistics
- Graph/Display Data

SEQUENCES MANIPULATION

- FASTA manipulation
- FastQC: fastq/sam/bam
- Illumina fastq
- Generic FASTQ manipulation
- FASTX-Toolkit for FASTQ data

SAM/BAM MANIPULATION : PICARD (BETA)

- Conversion
- QC/Metrics for sam/bam

### Upload File from Genotoul (Galaxy Version 1.0.0)

Options

**Path to file**

Path must be like : /work/USERNAME/somewhere/afile

**File type**

Execute

**What it does**

This program allows you to use a file stored in your genotoul work directory and optimize Galaxy work space by creating symlinks.

**Path to file**

This must be an absolute path to a file located in your genotoul work directory. The path must start with `/work/YOUR_USER_NAME/blablabla.extension`

valid path : `/work/LinuxUserName/galaxy/file.extension`  
invalid path : `/home/LinuxUserName/work/galaxy/file.extension`

**⚠ To use this tool and to maintain the confidentiality of yours directories:**

1. Create a "galaxy" directory in your work : `mkdir galaxy`
2. `chmod a+x /work/LinuxUserName`

Example : `drwxr-x--x 4 smaman sigenae 16384 mar 9 14:15 /work/smaman`- 3. `chmod a+r /work/LinuxUserName/dataGalaxy.fasta`

**ℹ Thanks to the fact that this tool requires you to enter your filepath (without "browse" button), you can manage your data privacy.**

For example, if your data to download in Galaxy are: `/work/LinuxUserName/galaxy/data.fasta`:

1. Add "x" rights to "others" on `/work/LinuxUserName/` and on `galaxy/`

It is not useful that "others" have "r" rights of these directories.- 2. Add "r" rights (only) to "others" on `data.fasta` file.

Thus, Galaxy can access and read `data.fasta` but all other files in those directories will not be accessible or readable.

**⚠ fastq files have to be uploaded in a correct format (for instance, `fatsqsanger`) in order to be used by Galaxy tools. If this is not the case, your fastq files uploaded will not be listed among available datasets in Galaxy tools.**

---

Version Galaxy Tool : V1.0

Versions of bioinformatics tools used : No bioinformatique tool used.

History

search datasets

merged

0 b

**ℹ** This history is empty. You can [load your own data](#) or [get data from an external source](#)

Copy the text here by pressing Ctrl+V.

Don't forget to change the Datatype!

**Path to file**

`/work/mleboulch/Formation`

Path must be like : `/work/USERNAME/somewhere/afile`



**Path to file**

`/work/mleboulch/Formation/100_10000seq_sample.tar.gz`

Path must be like : `/work/USERNAME/somewhere/afile`

You must add « / » and the name of the file at the end of the text.

Galaxy Analyze Data Workflow Shared Data Visualization Help User Using 813.2 MB

Tools ↑

search tools

MANAGE YOUR DATA FILES

**Get Data**

- [Upload File from your computer](#)
- [Upload File from Genotoul](#)
- [Upload several files from Genotoul without impact too much your Galaxy quota](#)
- [Upload ZIP file from an URL or browse your local files system](#)
- [UCSC Main table browser](#)
- [UCSC Test table browser](#)
- [UCSC Archaea table browser](#)
- [EBI SRA ENA SRA](#)

**Download Data**

**Jobs statistics**

FILES MANIPULATION

- [Text Manipulation](#)
- [Filter and Sort](#)
- [Join, Subtract and Group](#)
- [GFF](#)
- [BED Tools](#)
- [Convert Formats](#)
- [Fetch Sequences](#)
- [Statistics](#)
- [Graph/Display Data](#)

SEQUENCES MANIPULATION

- [FASTA manipulation](#)
- [FastQC: fastq/sam/bam](#)
- [Illumina fastq](#)
- [Generic FASTQ manipulation](#)
- [FASTX-Toolkit for FASTQ data](#)

SAM/BAM MANIPULATION : PICARD (BETA)

- [Conversion](#)
- [QC/Metrics for sam/bam](#)

History ↻ ⚙️ 📄

search datasets

**merged**

1 shown

4.55 MB ✔️ 🗑️ 💬

**1: /work/mleboulch /Formation /100\_10000seq\_sample.tar.gz** 👁️ ✎ ✖️

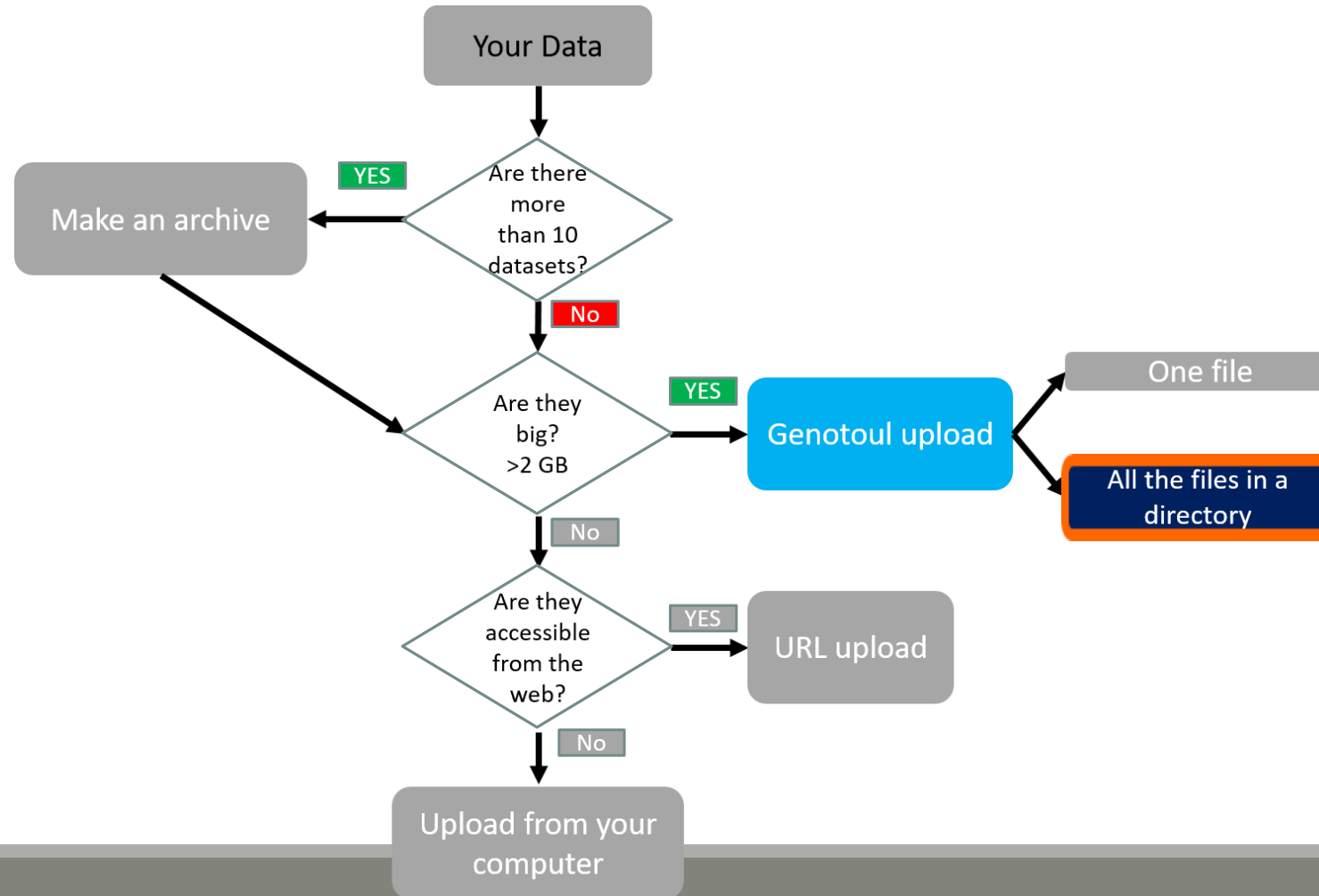
1 job has been successfully added to the queue - resulting in the following datasets:

**1: /work/mleboulch/Formation/100\_10000seq\_sample.tar.gz**

You can check the status of queued jobs and view the resulting data by refreshing the History pane. When the job has been run the status will change from 'running' to 'finished' if completed successfully or 'error' if problems were encountered.

After executing, the file is imported into Galaxy.

# Upload files from Genotoul



File Edit View Transfer Server Bookmarks Help

Host:  Username:  Password:  Port:  Quickconnect

Status: Setting permissions of /work/mleboulch/Formation/temp/temp2.fastq to 755  
 Status: /work/mleboulch/Formation/temp/temp2.fastq: 0644 -> 0755  
 Status: Retrieving directory listing of "/work/mleboulch/Formation"...  
 Status: Listing directory /work/mleboulch/Formation  
 Status: Directory listing of "/work/mleboulch/Formation" successful  
 Status: Retrieving directory listing of "/work/mleboulch/Formation/temp"...  
 Status: Listing directory /work/mleboulch/Formation/temp  
 Status: Directory listing of "/work/mleboulch/Formation/temp" successful

Local site: /home/malo/Bureau/Data/merged/

- Data
  - merged

Filename	Filesize	Filetype	Last modified
..			
100_10000seq_sample.tar.gz	4,773,706	gz-file	06/28/2017 05:...
100_10000seq_sampleC3.fastq	9,714,424	fastq-file	06/16/2015 10:...
100_10000seq_sampleC2.fastq	9,714,107	fastq-file	06/16/2015 10:...
100_10000seq_sampleC1.fastq	9,716,445	fastq-file	06/16/2015 10:...
100_10000seq_sampleB3.fastq	9,707,364	fastq-file	06/16/2015 10:...
100_10000seq_sampleB2.fastq	9,707,921	fastq-file	06/16/2015 10:...

10 files. Total size: 92,628,372 bytes

Remote site: /work/mleboulch/Formation

- Formation
  - temp

Filename	Filesize	Filetype	Last modified	Per
..				
100_10000seq_sample.tar.gz	4,773,706	gz-file	06/29/2017 06:04:35 PM	-rwx
temp		Directory	06/29/2017 06:01:32 PM	drwx

1 file and 1 directory. Total size: 4,773,706 bytes

Server/Local file | Direction | Remote file | Size | Priority | Status

Queued files | Failed transfers | Successful transfers (3)

Queue: empty

Double click on the temp directory.

Host:  Username:  Password:  Port:  Quickconnect 

Status: Listing directory /work/mleboulch/Formation  
Status: Directory listing of "/work/mleboulch/Formation" successful  
Status: Retrieving directory listing of "/work/mleboulch/Formation/temp"...  
Status: Listing directory /work/mleboulch/Formation/temp  
Status: Directory listing of "/work/mleboulch/Formation/temp" successful  
Status: Retrieving directory listing of "/work/mleboulch/Formation/temp"...  
Status: Listing directory /work/mleboulch/Formation/temp  
Status: Directory listing of "/work/mleboulch/Formation/temp" successful

Local site: /home/malo/Bureau/Data/merged/



Filename	Filesize	Filetype	Last modified
..			
100_10000seq_sample.tar.gz	4,773,706	gz-file	
100_10000seq_sampleC3.fastq	9,714,424	fastq-file	
100_10000seq_sampleC2.fastq	9,714,107	fastq-file	
100_10000seq_sampleC1.fastq	9,716,445	fastq-file	06/16/2015 10:...
100_10000seq_sampleB3.fastq	9,707,364	fastq-file	06/16/2015 10:...
100_10000seq_sampleB2.fastq	9,707,921	fastq-file	06/16/2015 10:...
100_10000seq_sampleB1.fastq	9,709,480	fastq-file	06/16/2015 10:...
100_10000seq_sampleA3.fastq	9,859,424	fastq-file	06/16/2015 10:...
100_10000seq_sampleA2.fastq	9,863,209	fastq-file	06/16/2015 10:...
100_10000seq_sampleA1.fastq	9,862,292	fastq-file	06/16/2015 10:...

10 files. Total size: 92,628,372 bytes

Remote site: /work/mleboulch/Formation/temp



Filename	Filesize	Filetype	Last modified	Per
..				
temp2.fastq	2,226,378	fastq-file	06/29/2017 06:01:33 PM	-rw
temp.fastq	2,226,378	fastq-file	06/29/2017 06:01:33 PM	-rw

2 files. Total size: 4,452,756 bytes



Copy this address.

Server/Local file	Direction	Remote file	Size	Priority	Status
-------------------	-----------	-------------	------	----------	--------

Queued files | Failed transfers | Successful transfers (3)

Queue: empty

Tools



search tools

MANAGE YOUR DATA FILES

**Get Data**[Upload File from your computer](#)[Upload File from Genotoul](#)[Upload several files from Genotoul without impact too much your Galaxy quota](#)[Upload ZIP file from an URL or browse your local files system](#)[UCSC Main table browser](#)[UCSC Test table browser](#)[UCSC Archaea table browser](#)[EBI SRA ENA SRA](#)[Download Data](#)[Jobs statistics](#)

FILES MANIPULATION

[Text Manipulation](#)[Filter and Sort](#)[Join, Subtract and Group](#)[GFF](#)[BED Tools](#)[Convert Formats](#)[Fetch Sequences](#)[Statistics](#)[Graph/Display Data](#)

SEQUENCES MANIPULATION

[FASTA manipulation](#)[FastQC: fastq/sam/bam](#)[Illumina fastq](#)[Generic FASTQ manipulation](#)[FASTX-Toolkit for FASTQ data](#)

SAM/BAM MANIPULATION : PICARD (BETA)

[Conversion](#)[QC/Metrics for sam/bam](#)

Upload several files from Genotoul without impact too much your Galaxy quota (Galaxy Version 1.0.1)

Options

Path to your directory which contains several files

Path must be like : /work/USERNAME/somewhere/

**What it does**

This program allows you to use a file stored in your genotoul work directory and optimize Galaxy work space by creating symlinks.

**Path to file**

This must be an absolute path to a file located in your genotoul work directory. The path must start with **/work/YOUR\_USER\_NAME/directory**

valid path : /work/LinuxUserName/directory

invalid path : /home/LinuxUserName/work/directory

**!** To use this tool and to maintain the confidentiality of yours directories:

1. Create a "galaxy" directory in your work : mkdir galaxy
2. chmod a+x /work/LinuxUserNam

Example : drwxr-x--x 4

**!** Thanks to the fact that this tool

For example, if your data to down

Add "x" rights to "others" on /work

It is not useful that "others" ha

Thus, Galaxy can access and rea

---

Version Galaxy Tool : V1.0

Versions of bioinformatics tools us

---

Contacts (noms et emails) : [sigenae](#)

E-learning available : Yes.

Please cite :

Depending on the help provided you can cite us in acknowledgements, references or both.

Examples : Acknowledgements We wish to thank the SIGENAE group for ....

References : SIGENAE [http://www.sigenae.org](#)

History



search datasets

temp

0 b

**!** This history is empty. You can [load your own data](#) or [get data from an external source](#)

- Go back to Galaxy and switch to temp history.
- Go to Get Data > Upload Several files from Genotoul
  - Paste the address into field.
  - Be careful all the files from the directory will be uploaded!

## Tools



search tools



## MANAGE YOUR DATA FILES

**Get Data**[Upload File](#) from your computer[Upload File](#) from Genotoul[Upload several files from Genotoul](#)  
without impact too much your  
Galaxy quota[Upload ZIP file](#) from an URL or  
browse your local files system[UCSC Main](#) table browser[UCSC Test](#) table browser[UCSC Archaea](#) table browser[EBI SRA](#) ENA SRA**Download Data****Jobs statistics**

## FILES MANIPULATION

[Text Manipulation](#)[Filter and Sort](#)[Join, Subtract and Group](#)[GFF](#)[BED Tools](#)[Convert Formats](#)[Fetch Sequences](#)[Statistics](#)[Graph/Display Data](#)

## SEQUENCES MANIPULATION

[FASTA manipulation](#)[FastQC: fastq/sam/bam](#)[Illumina fastq](#)[Generic FASTQ manipulation](#)[FASTX-Toolkit for FASTQ data](#)SAM/BAM MANIPULATION : PICARD  
(BETA)[Conversion](#)[QC/Metrics for sam/bam](#)

1 job has been successfully added to the queue - resulting in the following datasets:

**1: Upload several files from Genotoul**

You can check the status of queued jobs and view the resulting data by refreshing the History pane. When the job has been run the status will change from 'running' to 'finished' if completed successfully or 'error' if problems were encountered.

- Click on execute.

- All the files from the directory are uploaded.

## History



search datasets

**temp**

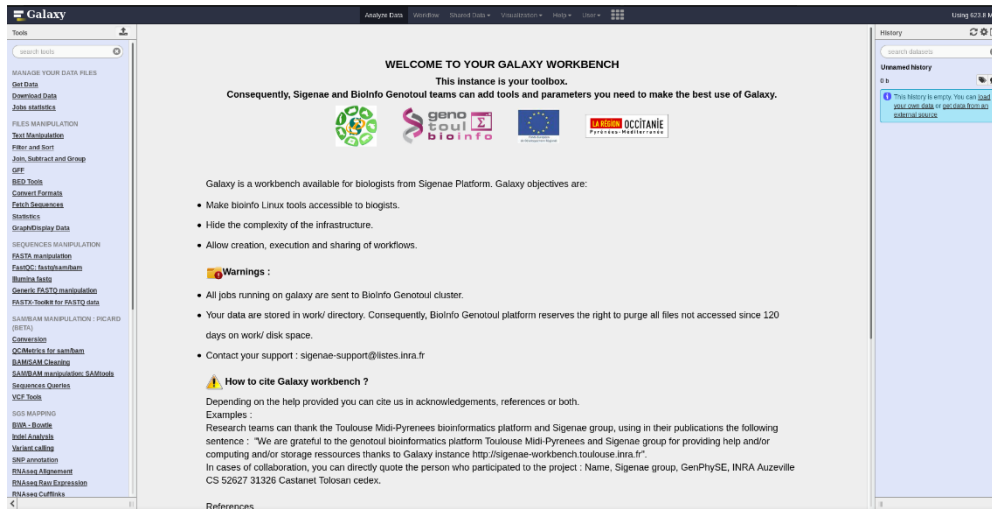
3 shown

4.25 MB

**3: Upload several files  
from Genotoul (temp2)****2: Upload several files  
from Genotoul (temp)****1: Upload several files  
from Genotoul**



# Upload by Genotoul



1TB of space



/work: 1TB (1024GB) of space  
/save: 250GB of space

- This method allows you to have more disk space and to upload bigger files.
- Files not used in the last 180 days can be purged at any time.

# Share a History

---

# Share a history

---

- You can share a history with another Galaxy user:
  - For working with your colleague.
  - For support, so we can help you better and faster.
- You can import shared history to your account too.

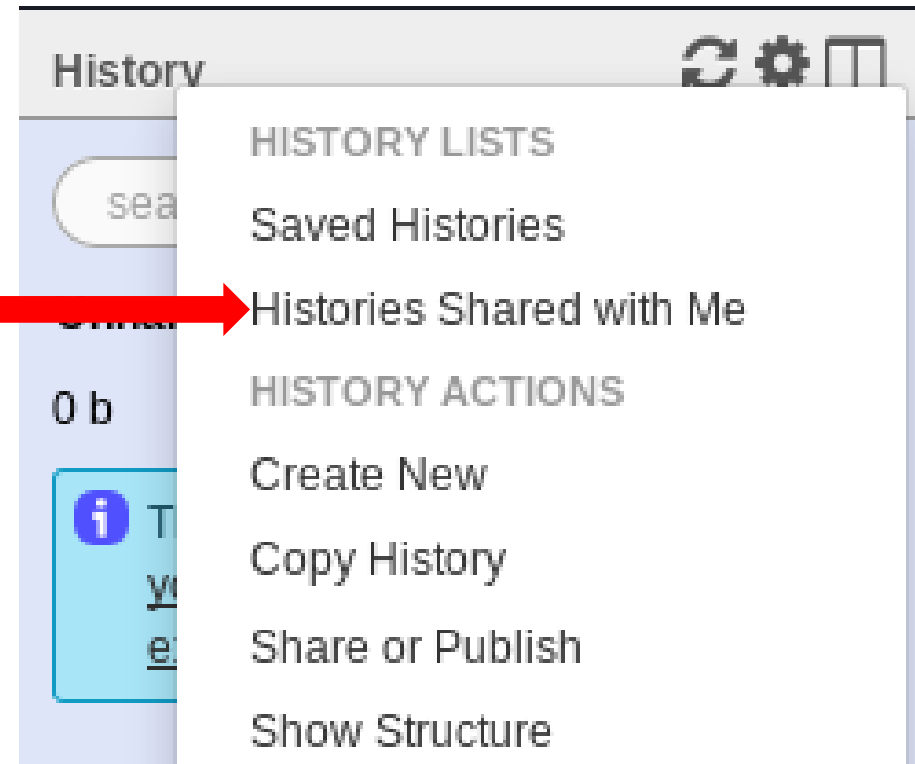
# Your Turn!

---

IMPORT A SHARED HISTORY TO YOUR ACCOUNT

# Import a shared history

Click on the wheel and click here.



Tools



## Histories shared with you by others

<input type="checkbox"/>	Name	Datasets	Created	Last Updated	Shared by
<input type="checkbox"/>	ASMA2016	27	Nov 18, 2016	Mar 09, 2017	geraldine.pascal@inra.fr
<input type="checkbox"/>	Copy of 'ASMA_2016_FE0802' shared by laurent.cauquil@toulouse.inra.fr (active items only)	24	Dec 09, 2016	Jan 03, 2017	geraldine.pascal@inra.fr

0 selected histories:

- Check the box and click on copy to import R1R2 history into your Galaxy account.
- You must import the « R1R2 » history.

Name of the history.

The person who shared the history.

History



search datasets

## Historique R1R2

3 shown

126.19 MB

**3: Upload several files from Genotoul (sampleA\_R2)****2: Upload several files from Genotoul (sampleA\_R1)****1: Upload several files from Genotoul**

empty

format: **txt**, database: ?

Epilog : job finished at Tue Jun 27 22:53:52 CEST 2017



# Your Turn!

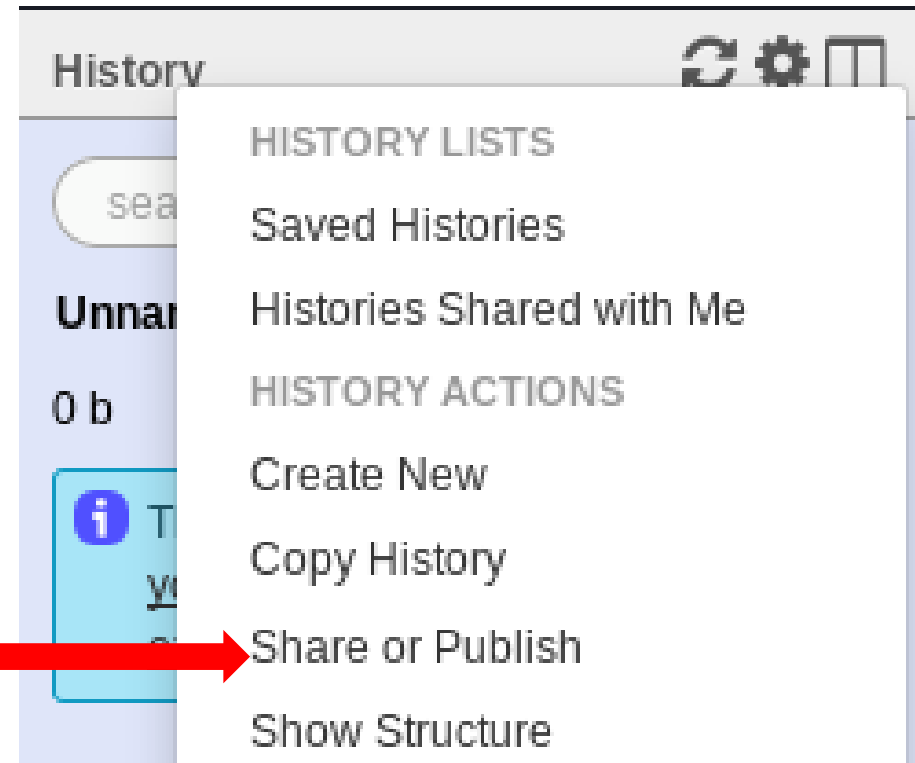
---

SHARE A HISTORY WITH YOUR NEIGHBOUR

# Share a history

---

- Switch to the history you want to share.
- Click on the wheel and click here.





Tools



search tools



## MANAGE YOUR DATA FILES

[Get Data](#)[Download Data](#)[Jobs statistics](#)

## FILES MANIPULATION

[Text Manipulation](#)[Filter and Sort](#)[Join, Subtract and Group](#)[GFF](#)[BED Tools](#)[Convert Formats](#)[Fetch Sequences](#)[Statistics](#)[Graph/Display Data](#)

## SEQUENCES MANIPULATION

[FASTA manipulation](#)[FastQC: fastq/sam/bam](#)[Illumina fastq](#)[Generic FASTQ manipulation](#)[FASTX-Toolkit for FASTQ data](#)SAM/BAM MANIPULATION : PICARD  
(BETA)[Conversion](#)[QC/Metrics for sam/bam](#)[BAM/SAM Cleaning](#)[SAM/BAM manipulation: SAMtools](#)[Sequences Queries](#)[VCF Tools](#)

## Share or Publish History 'Historique R1R2'

## Make History Accessible via Link and Publish It

This history is currently restricted so that only you and the users listed below can access it. You can:

[Make History Accessible via Link](#)

Generates a web link that you can share with other people so that they can view and import the history.

[Make History Accessible and Publish](#)

Makes the history accessible via link (see above) and publishes the history to Galaxy's [Published Histories](#) section, where it is publicly listed and searchable.

## Share History with Individual Users

You have not shared this history with any users.

[Share with a user](#)[Back to Histories List](#)

Click on « share with a user ».

History



search datasets



## Historique R1R2

3 shown

126.19 MB



**3: Upload several files from Genotoul (sampleA\_R2)**



**2: Upload several files from Genotoul (sampleA\_R1)**



**1: Upload several files from Genotoul**



empty

format: **txt**, database: ?

Epilog : job finished at Tue Jun 27  
22:53:52 CEST 2017



Tools



search tools



## MANAGE YOUR DATA FILES

[Get Data](#)[Download Data](#)[Jobs statistics](#)

## FILES MANIPULATION

[Text Manipulation](#)[Filter and Sort](#)[Join, Subtract and Group](#)[GFF](#)[BED Tools](#)[Convert Formats](#)[Fetch Sequences](#)[Statistics](#)[Graph/Display Data](#)

## SEQUENCES MANIPULATION

[FASTA manipulation](#)[FastQC: fastq/sam/bam](#)[Illumina fastq](#)[Generic FASTQ manipulation](#)[FASTX-Toolkit for FASTQ data](#)

## SAM/BAM MANIPULATION : PICARD (BETA)

[Conversion](#)[QC/Metrics for sam/bam](#)[BAM/SAM Cleaning](#)[SAM/BAM manipulation: SAMtools](#)[Sequences Queries](#)[VCF Tools](#)

Share 1 histories

## Histories to be shared:

History Name	Number of Datasets
Historique R1R2	3

## Galaxy user emails with which to share histories

Enter a Galaxy user email address or a comma-separated list of addresses if sharing with multiple users

- Enter an email address from a Galaxy user.
- Auto-completion helps you to find your co-worker

History



search datasets



## Historique R1R2

3 shown

126.19 MB

**3: Upload several files from Genotoul (sampleA\_R2)****2: Upload several files from Genotoul (sampleA\_R1)****1: Upload several files from Genotoul**

empty

format: **txt**, database: ?

EpiLog : job finished at Tue Jun 27 22:53:52 CEST 2017



## Galaxy user emails with which to share histories

malo.leboulch@inra.fr

malo.leboulch@inra.fr a-separat

Submit

- You must click on the address in the dropdown menu.
- Click on submit.
- The history will appear in your co-worker's history.

# Manipulate datasets

---

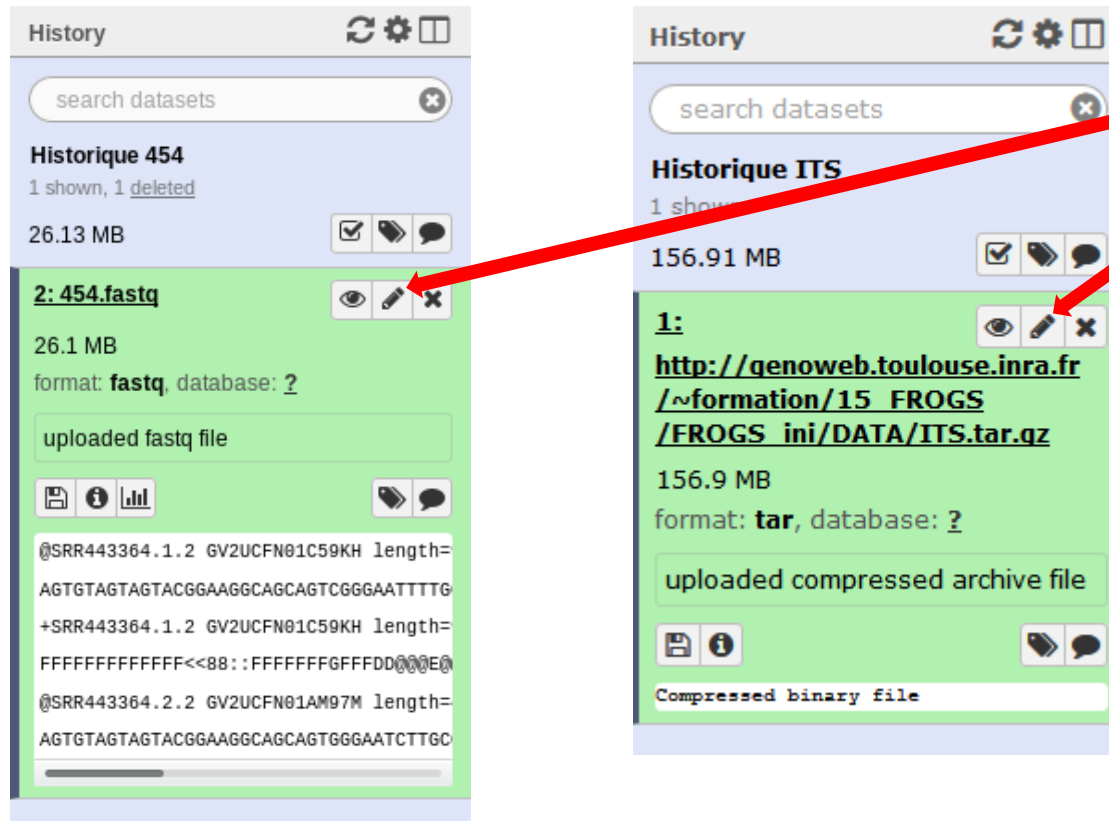
# Your Turn!

---

RENAME A DATASET

# Rename a dataset

- Switch to 454 history or ITS history



Click here to display attributes and change the name.

# Rename a dataset

Attributes Convert Format Datatype Permissions

Edit Attributes

**Name:**  
http://genoweb.toulouse.inra.fr/~formatio

**Info:**  
uploaded fastq file

**Annotation / Notes:**

Add an annotation or notes to a dataset; annotations are available when a history is viewed.

**Database/Build:**  
unspecified (?)

Save

Auto-detect

This will inspect the dataset and attempt to correct the above column values if they are not accurate.

OR

Attributes Convert Format Datatype Permissions

Edit Attributes

**Name:**  
\_FROGS/FROGS\_ini/DATA/ITS.tar.gz

**Info:**  
uploaded compressed archive file

**Annotation / Notes:**

Add an annotation or notes to a dataset; annotations are available when a history is viewed.

**Database/Build:**  
unspecified (?)

Save

Auto-detect

This will inspect the dataset and attempt to correct the above column values if they are not accurate.

# Rename a dataset

Attributes Convert Format Datatype Permissions

Edit Attributes

**Name:**

**Info:**

**Annotation / Notes:**

Add an annotation or notes to a dataset; annotations are available when a history is viewed.

**Database/Build:**

Save

Auto-detect

This will inspect the dataset and attempt to correct the above column values if they are not accurate.

If you put the wrong datatype in during the upload, you can change it here.



# Rename a dataset

---

- Do the same with the merged history:
  - Switch to the merged history.
  - Change the name of the file to «100\_10000seq\_sample.tar.gz ».

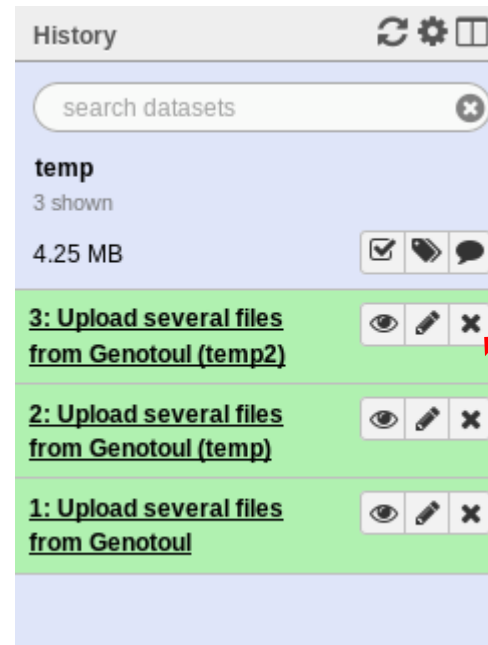
# Your Turn!

---

DELETE A DATASET

# Delete a dataset

- Switch to temp history.



Click here to delete a dataset.

# Delete a dataset

The image shows two screenshots of the Galaxy History interface. The left screenshot shows a dataset named 'temp' with a size of 4.25 MB and a status of '1 deleted'. A blue callout box with the text 'Click here.' has a red arrow pointing to the '1 deleted' link. Below the dataset information are two entries: '2: Upload several files from Genotoul (temp)' and '1: Upload several files from Genotoul'. A large grey arrow points from the left screenshot to the right screenshot. The right screenshot shows the same dataset 'temp' with a size of 4.25 MB and a status of '3 shown, hide deleted'. A yellow warning box is displayed over the dataset, containing the text 'This dataset has been deleted', 'Undelete it', and 'Permanently remove it from disk'. A red arrow points from a blue callout box with the text 'Delete this dataset permanently.' to the 'Permanently remove it from disk' link. Below the dataset information are three entries: '3: Upload several files from Genotoul (temp2)', '2: Upload several files from Genotoul (temp)', and '1: Upload several files from Genotoul'.

Click here.

temp  
2 shown, 1 deleted  
4.25 MB

2: Upload several files from Genotoul (temp)

1: Upload several files from Genotoul

History

search datasets

temp  
3 shown, [hide deleted](#)  
4.25 MB

This dataset has been deleted  
Undelete it  
[Permanently remove it from disk](#)

3: Upload several files from Genotoul (temp2)

2: Upload several files from Genotoul (temp)

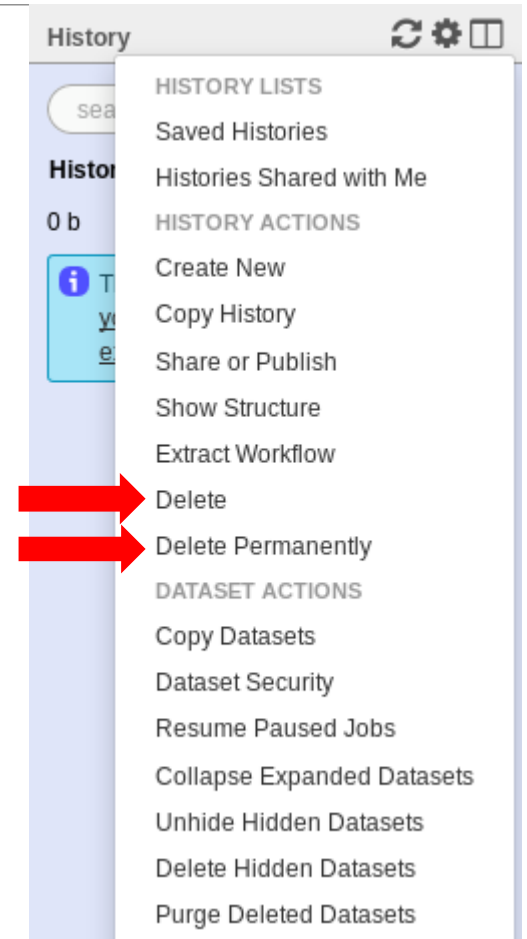
1: Upload several files from Genotoul

Delete this dataset permanently.

Nothing is deleted in Galaxy until you delete it permanently (=purge).

# How to delete a history?

- Stay in the temp history.
- Click on the wheel.
- Click on delete.
- A deleted item on Galaxy is recoverable.
- To definitively delete it:  
click on « Delete Permanently ».



Current History



Switch to



Switch to



Switch to



**multiplex**

2 shown, 1 [deleted](#)

2.12 MB



search datasets

*Drag datasets here to copy them to the current history*

**2: multiplex.fastq**



**1: barcode\_forward.tabular**



**merged**

1 shown

4.55 MB



search datasets

**1:**



**100\_10000seq\_sample.tar.gz**

**454**

1 shown

26.13 MB



search datasets

**1: 454.fastq**



**Historique R1R2**

2 shown, 1 [deleted](#)

126.19 MB



search datasets

**3: sampleA\_R2**



**2: sampleA\_R1**



# One word about data collections

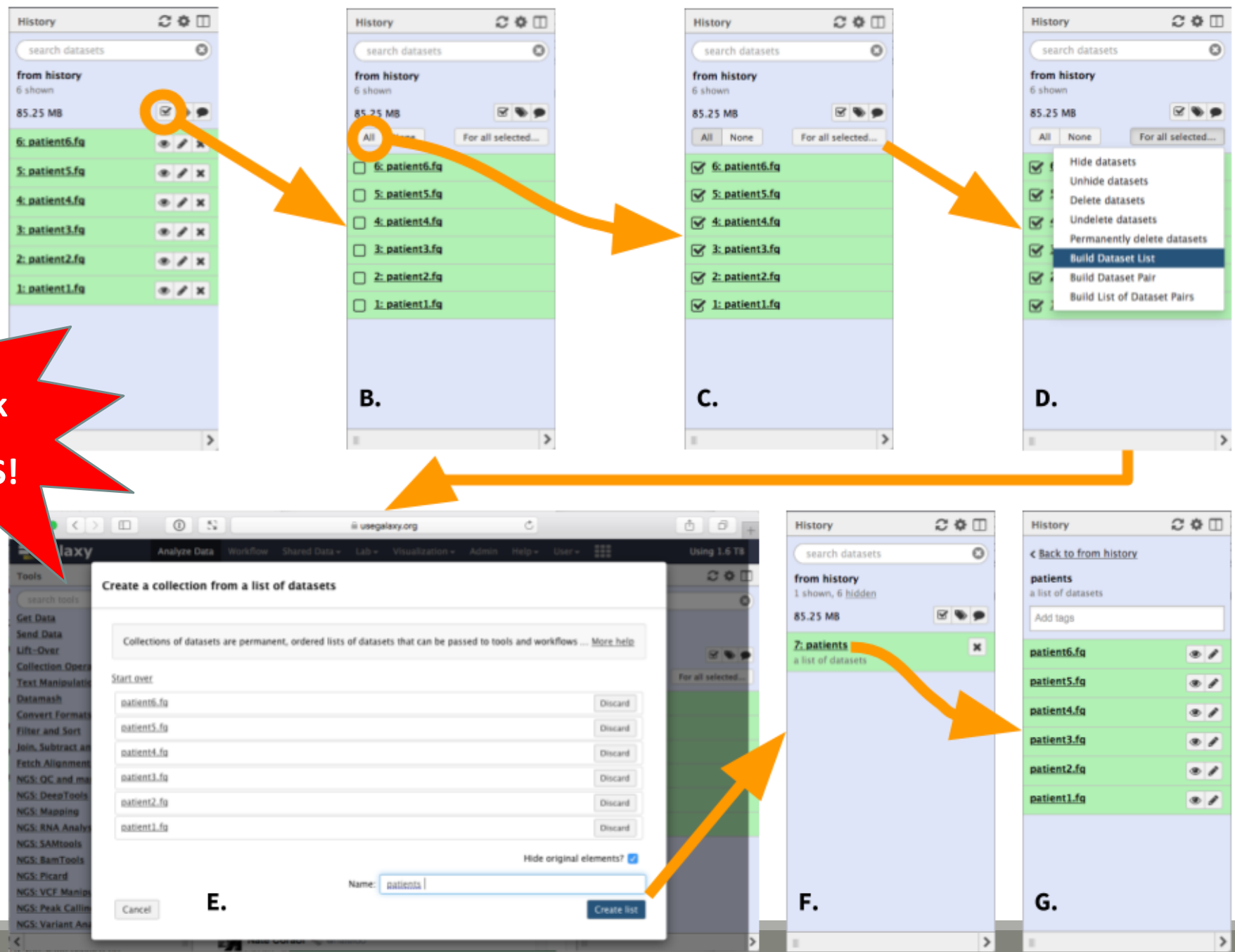
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- Used to perform the same analysis on multiple files.
- Gather multiple datasets in one collection.



**Don't work with FROGS!**

**Don't work  
with FROGS!**





# Others tools available on galaxy

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- Text Manipulation
- Filter and Sort
- FASTA manipulation

# Galaxy support

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- Mail: [support.sigenae@inra.fr](mailto:support.sigenae@inra.fr)
- If you need more training about bioinformatics and Galaxy, please connect to Sigenae e-learning platform: <http://sig-learning.toulouse.inra.fr/>

# How to cite Galaxy?

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- Goecks, J, Nekrutenko, A, Taylor, J and The Galaxy Team. "*Galaxy: a comprehensive approach for supporting accessible, reproducible, and transparent computational research in the life sciences.*" Genome Biol. 2010 Aug 25;11(8):R86.
- Blankenberg D, Von Kuster G, Coraor N, Ananda G, Lazarus R, Mangan M, Nekrutenko A, Taylor J. "*Galaxy: a web-based genome analysis tool for experimentalists*". Current Protocols in Molecular Biology. 2010 Jan; Chapter 19:Unit 19.10.1-21.
- Giardine B, Riemer C, Hardison RC, Burhans R, Elnitski L, Shah P, Zhang Y, Blankenberg D, Albert I, Taylor J, Miller W, Kent WJ, Nekrutenko A. "*Galaxy: a platform for interactive large-scale genome analysis.*" Genome Research. 2005 Oct; 15(10):1451-5.

# How to cite Genotoul Galaxy workbench?

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- Research teams can thank the Toulouse Midi-Pyrenees bioinformatics platform and Sigenae group, using in their publications the following sentence : "We are grateful to the genotoul bioinformatics platform Toulouse Midi-Pyrenees and Sigenae group for providing help and/or computing and/or storage resources thanks to Galaxy instance <http://sigenae-workbench.toulouse.inra.fr>".
- In cases of collaboration, you can directly quote the person who participated to the project : Name, Sigenae group, GenPhySE, INRA Auzeville CS 52627 31326 Castanet Tolosan cedex.