

A - Training on Galaxy: Metabarcoding

October 2024 - webinar

GALAXY Practice

LUCAS AUER, MARIA BERNARD, LAURENT CAUQUIL, VINCENT DARBOT, MAHENDRA MARIADASSOU, GÉRALDINE PASCAL & OLIVIER RUÉ













Objectives of the 1st part

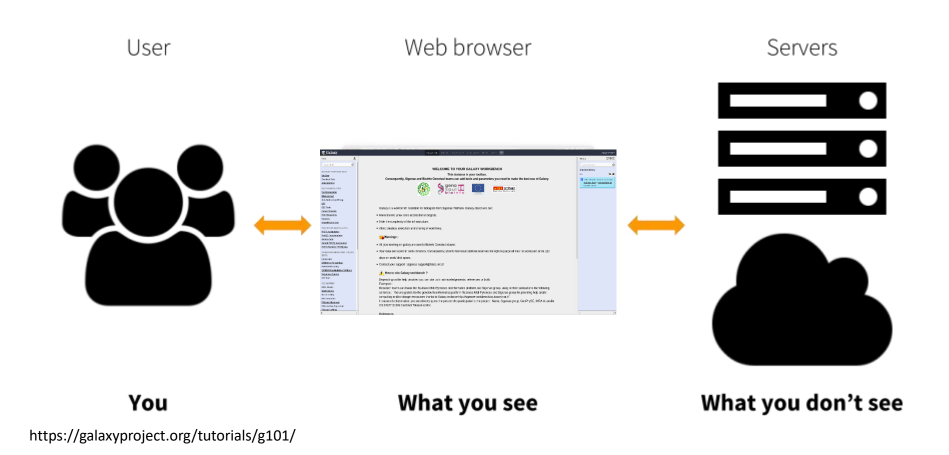
- Learn the basics of Galaxy
- Prepare the 1 dataset for FROGS formation

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



Where to use Galaxy?

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





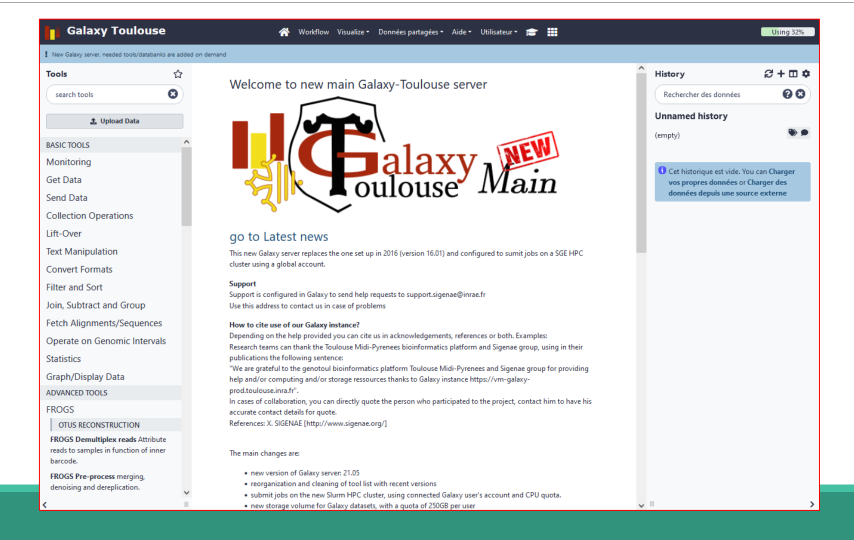
Where to use Galaxy?

- The Toulouse Galaxy platform: https://vm-galaxy-prod.toulouse.inrae.fr
- Institut Français Bioinformatique IFB (national server): https://metabarcoding.usegalaxy.fr
- Migale Plateform: galaxy.migale.inra.fr
- Others...
- ↑ The different platforms of Galaxy are not connected together
- Galaxy is installed on many clusters across the world.
- Some tools are in <u>our</u> Galaxy platform but not in <u>other</u> platforms.
- Your data is not shared with other Galaxy platforms than ours.

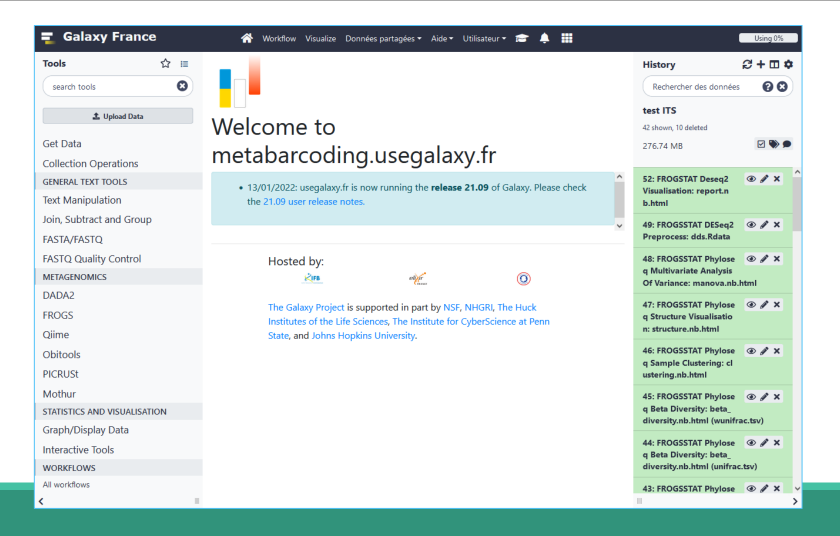
Exemple of Galaxy platforms



Exemple of Galaxy platforms



Exemple of Galaxy platforms



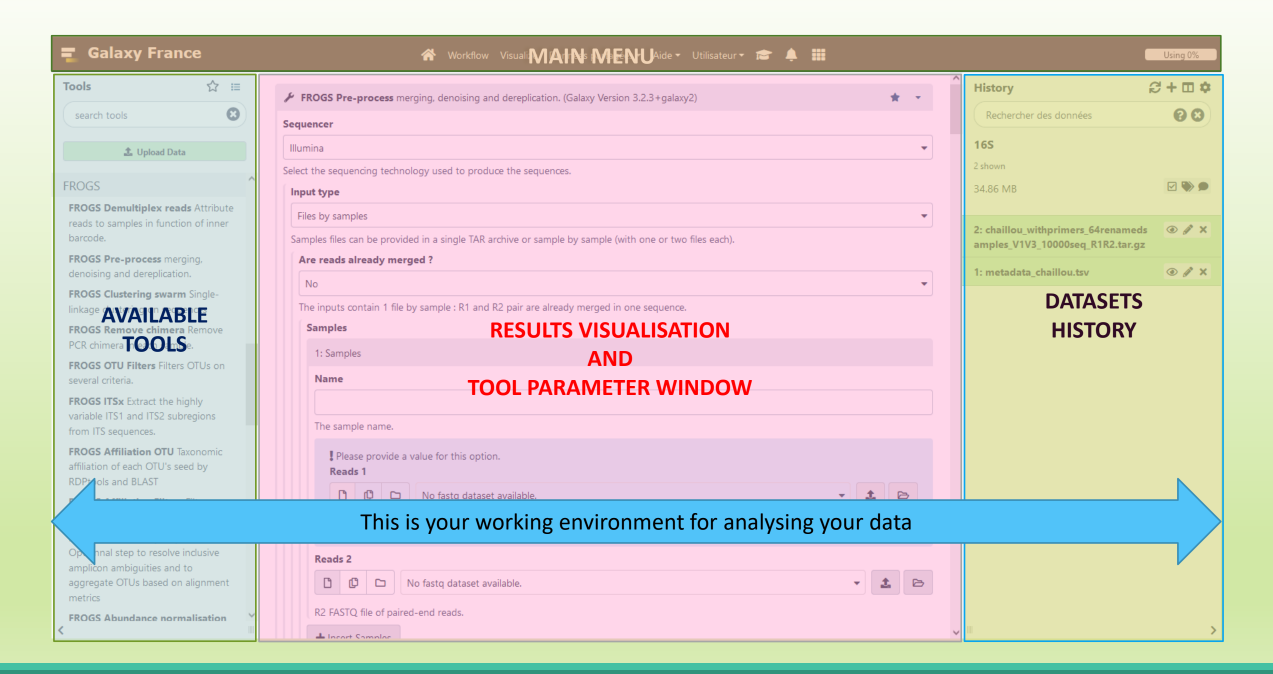
Practice:

CONNECT TO GALAXY WORKBENCH

Exercise

- Go to https://metabarcoding.usegalaxy.fr
- Enter your login and your password

Welcome to Galaxy, please log in
Public Name or Email Address
Password
Forgot password? Click here to reset your password. Login
Don't have an account? Registration for this Galaxy instance is disabled. Please contact an administrator for assistance.



Vocabulary of Galaxy

Tools:

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

Datasets:

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

Histories:

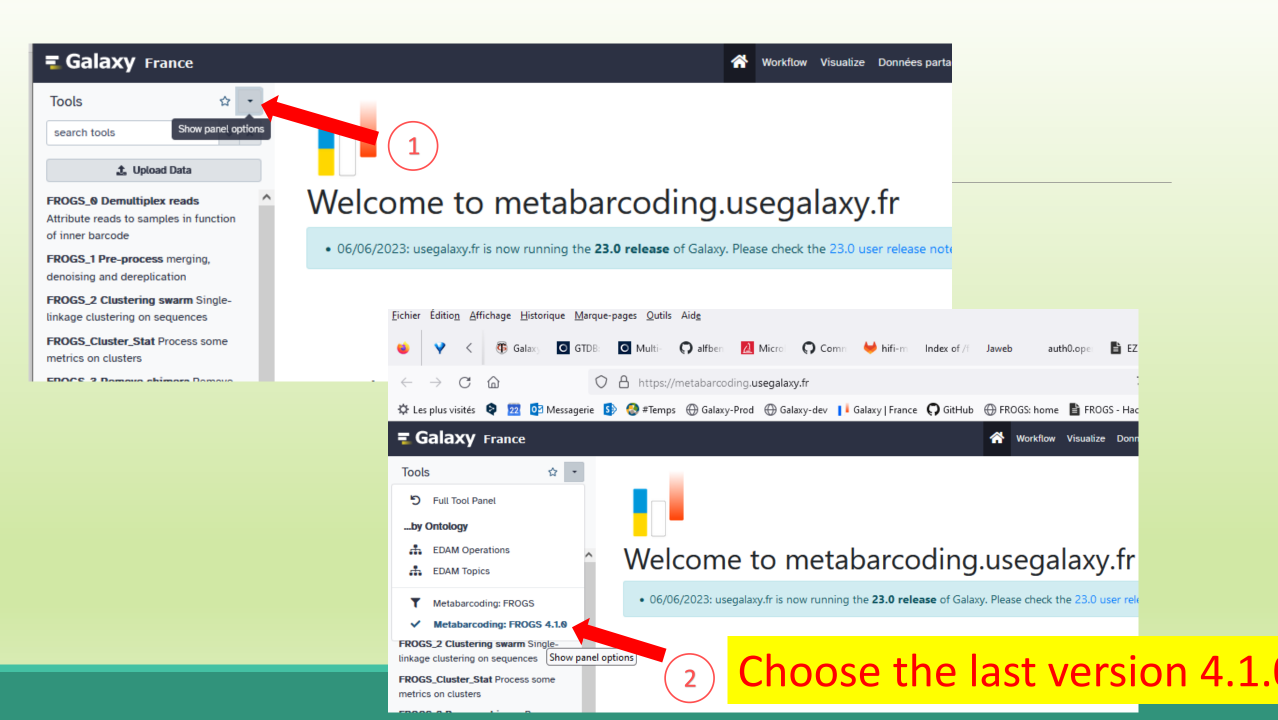
- A tool generates datasets and these datasets are stored in the current history.
- Everything is permanently saved.
- If you log off your computer or browser, it's ok, everything will keep running and be saved!

Pratice:

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.



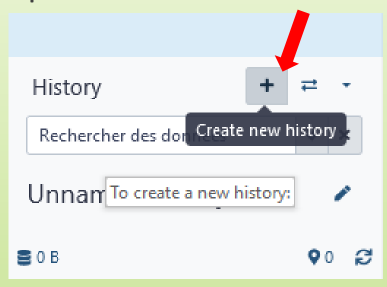
Manipulate Histories

Practice:

CREATE THE NECESSARY HISTORY FOR FROGS TRAINING

To create a new history:

Click on the plus.



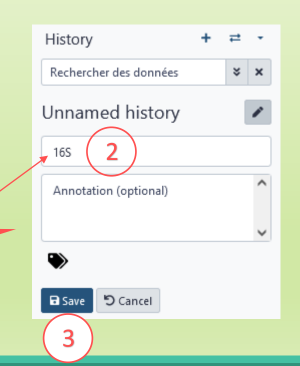
To rename a history:

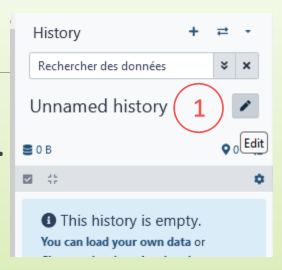
Click on the pen next to "Unnamed history" (top right).



Hit « Enter » to validate.

Don't use special characters or accents!

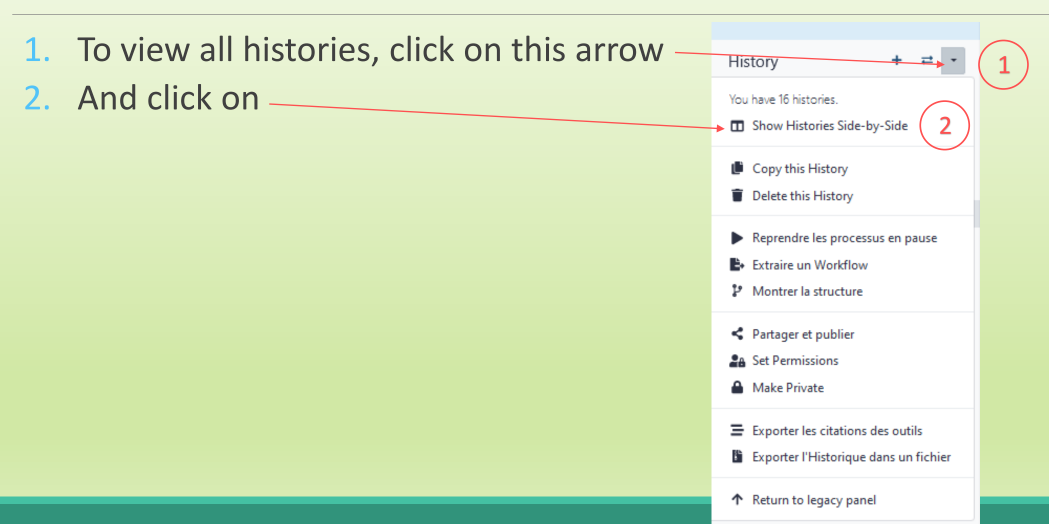




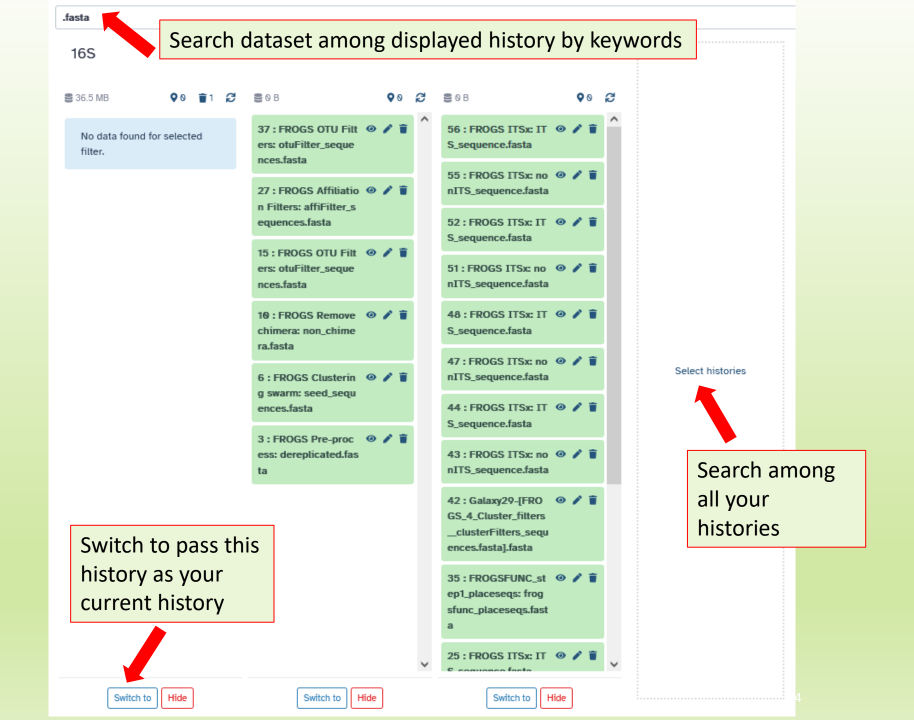
Exercise

 Create another history as before with 16S. This new history will be named ITS

How to list all histories?



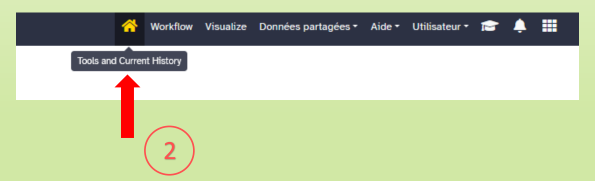
Explore the « View all histories » section



Switch current history

1. When you are several histories, switch to the history named « 16S » as current history

2. You can click on "home" to go back to the main interface.





Data import

How to import your data to Galaxy?

- Several ways to upload your data to Galaxy:
 - From your computer



By URL



Shared by other users of Galaxy

drag-and-drop files

Paste an url



Formation data

All the training data are online on:
 https://web-genobioinfo.toulouse.inrae.fr/~formation/15 FROGS/Webinar data/

Index of /~formation/15_FROGS/Webinar_data



Remark: how create an archive?

Not for this session, but for your own data, it will be important to know how create an archive.

To learn how create an archive

https://web-genobioinfo.toulouse.inrae.fr/~formation/15 FROGS/Webinar data/How create an archive.pdf

Create an archive is always possible!

It is our recommendation.

Practice: 16S history

UPLOAD FILE FROM AN URL

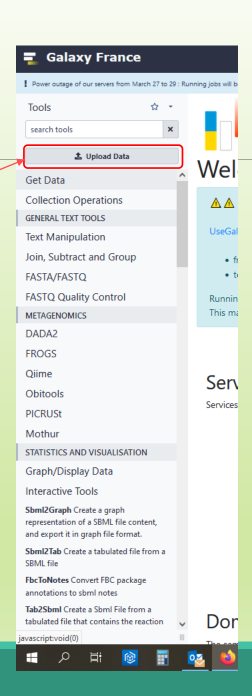
How to import your data to Galaxy?

Several ways to upload your data to Galaxy:



16S history creation

- In your current history "165" (If it not the case, switch to 16S history)
- 2. Go to **Upload Data** on the left panel



16S history creation

Parent Directory How create an archive.pdf

Apache/2.2.15 (CentOS

Inspecter

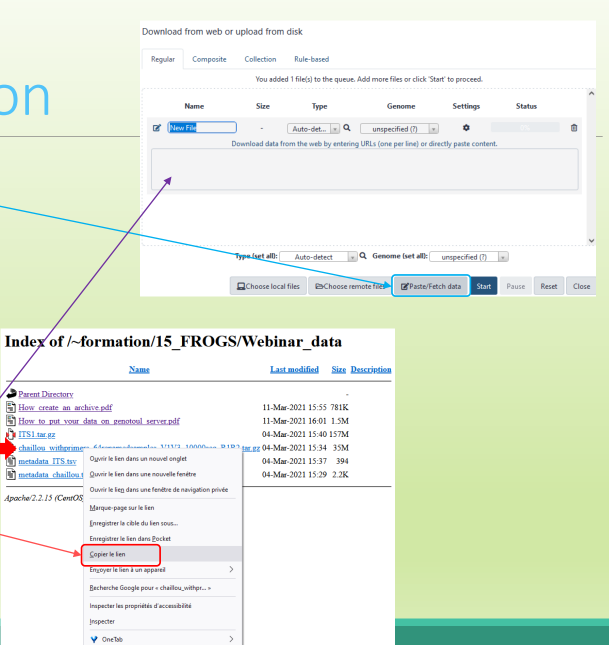
Video DownloadHelper

Click on Paste/Fetch Data

Collect your data address on https://webgenobioinfo.toulouse.inrae.fr/~formation/1 5 FROGS/Webinar data/

Click right on the name file

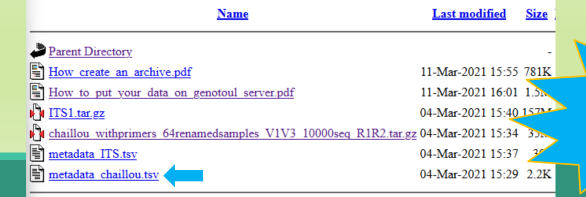
- Copy the address of the link
- Paste the link in the grey window

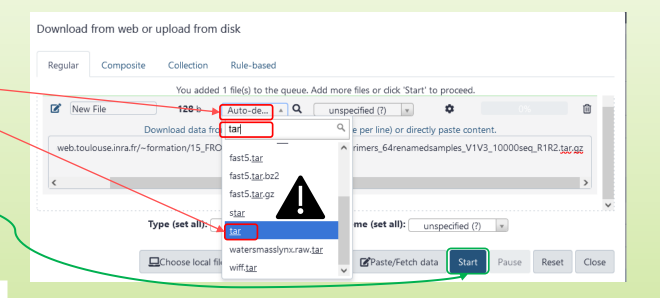


16S history creation

- 7. Change the data type!
- 8. Click on Start
- Start again with the file metadata_chaillou.tsv

Index of /~formation/15_FROGS/Webinar_data





Remember to change your data type to TSV

Share a History

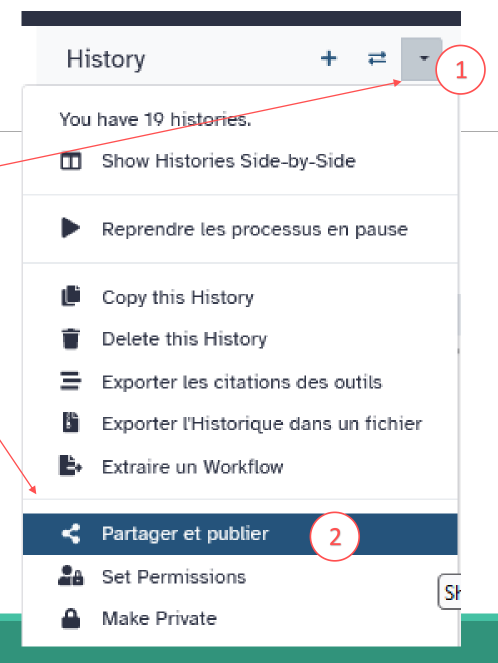
Share a history

- 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.

Share a history

To share an history

- 1. Click on this arrow
- 2. Click on share and publish



Share a history

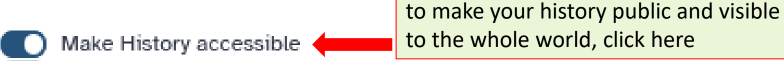
Share or Publish History 'test frogs sans clustering'

Make History accessible Access to this History is currently restricted so that only you and the users listed below can access it. Note that sharing a History will also allow access to l of its To share click on and enter a user Share History with Individual Users -The following users will see this History in their History list and will be able to view, import and run it. lucas.auer@inrae.fr × Cancel Save To unshare click on the cross



Share a history

Share or Publish History 'test frogs sans clustering'





This History is currently accessible via link.

Anyone can view and import this History by visiting the following URL:

url: https://metabarcoding.usegalaxy.fr/u/geraldine.pascal/h/test-frogs-sans-clustering

Practice: share a history

SHARE 16S AND ITS HISTORIES WITH TRAINERS

Share your data with us ***



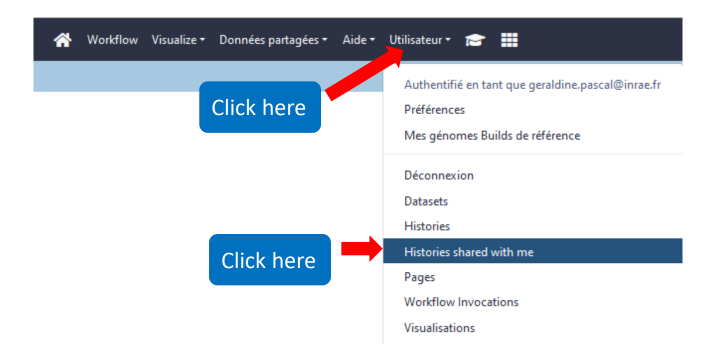
- When your history "16S" is ready and contains the 2 files
 - chaillou_withprimers_64renamedsamples_V1V3_10000seq_R1R2.tar.gz
 - metadata_chaillou.tsv
- Then share with

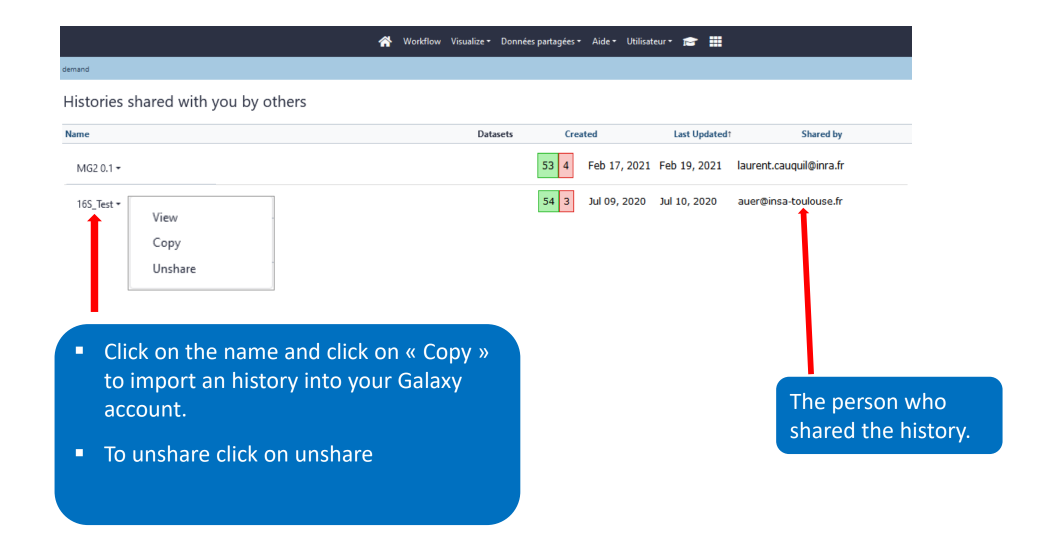
lucas.auer@inrae.fr × geraldine.pascal@inrae.fr ×

- Start again with the "ITS" history
- Contact us by email frogs-training@inrae.fr when it is done.

Other useful things

Import a shared history





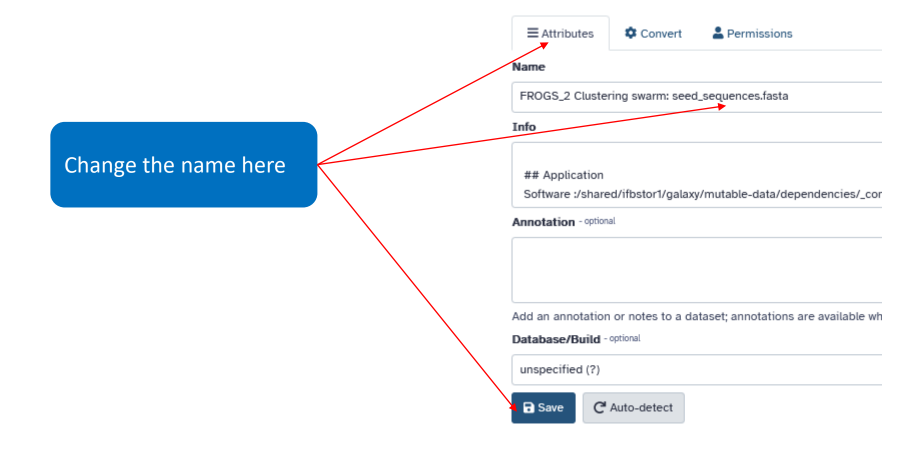
How manipulate datasets

To rename a dataset

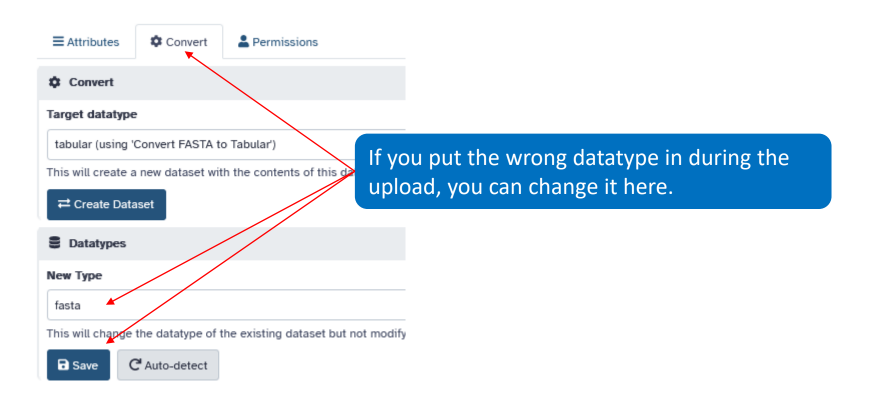


Click here to display attributes and change the name.

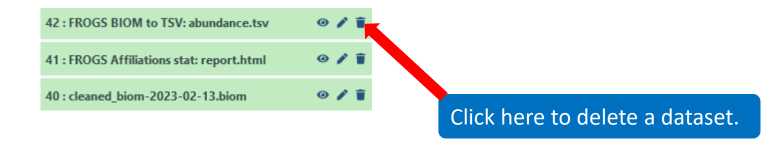
To rename a dataset



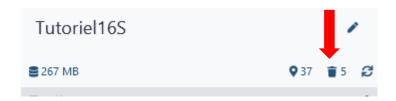
To change the datatype



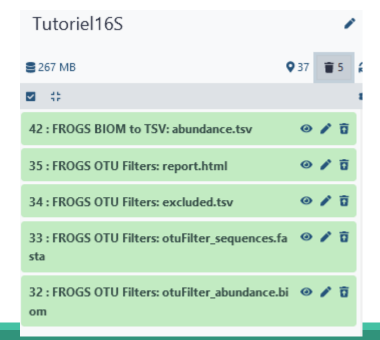
To delete a dataset



To see deleted dataset

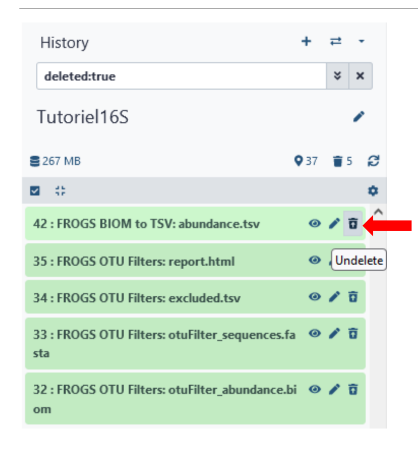


Click here to see deleted dataset.



Here the complete list of deleted dataset.

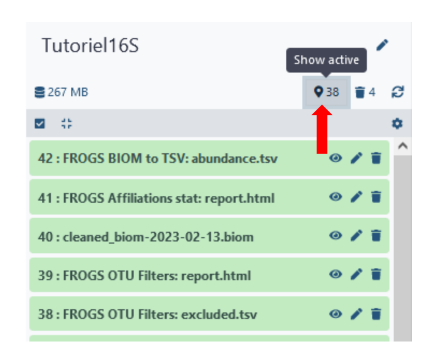
To undelete dataset



Click here and this file return into the history

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

To undelete dataset



Click here to sea again your active history

How to cite Galaxy workbench?

For Genotoul Bioinfo:

- Research teams can thank the Toulouse Occitanie bioinformatics platform and Sigenae group, using in their publications the following sentence: "We are grateful to the genotoul bioinformatics platform Toulouse Occitanie (Bioinfo Genotoul, doi: 10.15454/1.5572369328961167E12) and Sigenae group for providing help and/or computing and/or storage ressources 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, INRAE Auzeville CS 52627 31326 Castanet Tolosan cedex.

For IFB:

• We are grateful to the Institut Français de Bioinformatique (ANR-11-INBS-0013) for providing help and/or computing and/or storage resources.

For Migale:

- Research teams that have used Migale services are expected to thank the Migale bioinformatics facility in the publications of their analyses with the following sentence: "We are grateful to the INRAE MIGALE bioinformatics facility (MIGALE, INRAE, 2020. Migale bioinformatics Facility, doi: 10.15454/1.5572390655343293E12) for providing help and/or computing and/or storage resources"
- In cases of collaboration, you can directly quote the person who participated to the project with the two following affiliations: Université Paris-Saclay, INRAE, MalAGE, 78350, Jouy-en-Josas, France Université Paris-Saclay, INRAE, BioinfOmics, MIGALE bioinformatics facility, 78350, Jouy-en-Josas, France

Thank you for following this first step, see you on Monday 7 October!

YOUR TRAINERS: GÉRALDINE & LUCAS



