

From NCBI's Sequence Read Archive (SRA) to Galaxy: SARS-CoV-2 variant analysis

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Overview

? Questions

• Learn how to get and use data from the Sequence Read Archive in Galaxy.

Objectives

- Understand how Galaxy and the Sequence Read Archive interact.
- Be able to go from Galaxy to the Short Reach Archive, query SRA, use the SRA Run Selector to send selected metadata to Galaxy, and then import sequence data from SRA into Galaxy.

Requirements

- Introduction to Galaxy Analyses
- Sequence analysis
 - Quality Control: slides hands-on
 - Mapping: slides hands-on

▼ Time estimation: 45 minutes

☑ Supporting Materials

Last modification: Feb 12, 2021

Introduction

Now what?

Variation Analysis of SARS-Cov-2 sequencing

Introduction

The aim of this tutorial is to introduce you to the processing of next generation sequencing data in Galaxy. This tutorial uses a COVID-19 variant calling from Illumina data, but it isn't about variant calling *per se*.

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data

Get the reference genome data

Conclusion

Feedback

Citing this Tutorial At the completion of this tutorial you will know:

- How to find data in SRA and transfer this information to Galaxy
- How to perform basic NGS data processing in Galaxy including:
 - Quality Control (QC) of Illumina data
 - Mapping
 - Removal of duplicates
 - Variant calling with lofreq
 - Variant annotation
- Using datasets collections
- Importing data to Jupyter

Find necessary data in SRA

First we need to find a good dataset to play with. The Sequence Read Archive (SRA) is the primary archive of *unassembled reads* operated by the US National Institutes of Health (NIH). SRA is a great place to get the sequencing data that underlie publications and studies. Let's do that:

Hands-on: Task description

- 1. Go to NCBI's SRA page by pointing your browser to https://www.ncbi.nlm.nih.gov/sra
- 2. In the search box enter SARS-CoV-2 Patient Sequencing From
 Partners / MGH: Find data (Alternatively, you simply click on this
 link)
- The web page will show a large number of SRA datasets (at the time of writing there were 2,223). This is data from a study describing analysis of SARS-CoV-2 in Boston area.
- 4. Download metadata describing these datasets by:
 - clicking on Send to: dropdown
 - Selecting File
 - Changing Format to RunInfo
 - Clicking Create file Here is how it should look like:



5. This would create a rather large SraRunInfo.csv file in your Downloads folder.

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Now that we have downloaded this file we can go to a Galaxy instance and start processing it.

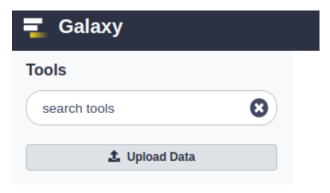
Comment :

Note that the file we just downloaded is **not** sequencing data itself. Rather, it is *metadata* describing properties of sequencing reads. We will filter this list down to just a few accessions that will be used in the remainder of this tutorial.

Process and filter SraRunInfo.csv file in Galaxy

Hands-on: Upload SraRunInfo.csv file into Galaxy

- Go to your Galaxy instance of choice such as one of the usegalaxy.org, usegalaxy.eu, usegalaxy.org.au or any other. (This tutorial uses usegalaxy.org).
- 2. Click Upload Data button:



- 3. In the dialog box that would appear click "Choose local files" button: Choose local
- 4. Find and select SraRunInfo.csv file from your computer
- 5. Click Start button
- 6. Close dialog by pressing **Close** button
- 7. You can now look at the content of this file by clicking (eye) icon. You will see that this file contains a lot of information about individual SRA accessions. In this study every accession corresponds to an individual patient whose samples were sequenced.

Galaxy can process all 2,000+ datasets but to make this tutorir'' need to selected a smaller subset. In particular our previous expression of the control of

data shows two interesting datasets SRR11954102 and SRR12733957 . So, let's pull them out.

Beware of Cuts

The Hands-on section below uses **Cut** tool. There are two **cut** tools in Galaxy due to historical reasons. This example uses tool with the full name **Cut columns from a table (cut)**. However, the same logic applies to the other tool. It simply has a slightly different interface.

Hands-on: Creating a subset of data

1. Find "Select lines that match an expression" tool in Filter and Sort section of the tool panel.

Ō	Tip:	Finding	tools	•
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- 2. Make sure the SraRunInfo.csv dataset we just uploaded is listed in the "Select lines from" field of the tool form.
- 3. In "the pattern" field enter the following expression → SRR12733957 | SRR11954102 . These are two accession we want to find separated by the pipe symbol | . The | means or : find lines containing SRR12733957 or SRR11954102 .
- 4. Click Execute button.
- 5. This will generate a file containing two lines (well ... one line is also used as the header, so it will appear the the file has three lines. It is OK.)
- 6. Cut the first column from the file using **/** "Cut" tool, which you will find in **Text Manipulation** section of the tool pane.
- 7. Make sure the dataset produced by the previous step is selected in the "File to cut" field of the tool form.
- 8. Change "Delimited by" to Comma
- 9. In "List of fields" select Column: 1.
- 10. Hit Execute This will produce a text file with just two lines:

SRR12733957 SRR11954102

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Now that we have identifiers of datasets we want we need to download the actual sequencing data.

Download sequencing data with Faster Download and Extract Reads in FASTQ

Hands-on: Task description

- Faster Download and Extract Reads in FASTQ with the following parameters:
 - o "select input type": List of SRA accession, one per line
 - The parameter [] "sra accession list" should point the output of the "Cut" from the previous step.
 - Click the Execute button. This will run the tool, which retrieves
 the sequence read datasets for the runs that were listed in the
 SRA dataset. It may take some time. So this may be a good
 time to do get coffee.
- 2. Several entries are created in your history panel when you submit this job:
 - Pair-end data (fasterq-dump) : Contains Paired-end datasets (if available)
 - Single-end data (fasterq-dump) Contains Single-end datasets (if available)
 - Other data (fasterq-dump) Contains Unpaired datasets (if available)
 - fasterq-dump log Contains Information about the tool execution

The first three items are actually *collections* of datasets. *Collections* in Galaxy are logical groupings of datasets that reflect the semantic relationships between them in the experiment / analysis. In this case the tool creates a separate collection each for paired-end reads, single reads, and *other*. See the Collections tutorials for more.

Explore the collections by first **clicking** on the collection name in the history panel. This takes you inside the collection and shows you the datasets in it. You can then navigate back to the outer level of your history.

Once fasterg finishes transferring data (all boxes are green

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ready to analyze it.

Now what?

You can now analyze the retrieved data using any sequence analysis tools and workflows in Galaxy. SRA holds backing data for every imaginable type of *-seq experiment.

If you ran this tutorial, but retrieved datasets that you were interested in, then see the rest of the GTN library for ideas on how to analyze in Galaxy.

However, if you retrieved the datasets used in this tutorial's examples above, then you are ready to run the SARS-CoV-2 variant analysis below.

Variation Analysis of SARS-Cov-2 sequencing data

In this part of the tutorial we will perform variant calling and basic analysis of the datasets downloaded above. We will start by downloading the Wuhan-Hu-1 SARS-CoV-2 reference sequence, then run adapter trimming, alignment and variant calling and finally look at the geographic distribution of some of the found variants.

The usegalaxy.* COVID-19 analysis project

This tutorial uses a subset of the data and runs through the Variation Analysis section of covid19.galaxyproject.org. The data for covid19.galaxyproject.org is being updated continuously as new datasets are made public.

Get the reference genome data

The reference genome data for today is for SARS-CoV-2, "Severe acute respiratory syndrome coronavirus 2 isolate Wuhan-Hu-1, complete genome", having the accession ID of NC_045512.2.

This data is available from Zenodo using the following link.

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Hands-on: Get the reference genome data

1. Import the following file into your history:

https://ftp.ncbi.nlm.nih.gov/genomes/all/GCF/009/858/895/GCF_009858895.2_ASM985889v3_genomic.fna.gz

Adapter trimming with fastp

Removing sequencing adapters improves alignments and variant calling. **fastp** \nearrow can automatically detect widely used sequencing adapters.

Hands-on: Task description

- 1. **fastp /** with the following parameters:
 - o "Single-end or paired reads": Paired Collection
 - [] "Select paired collection(s)": list_paired (output of Faster Download and Extract Reads in FASTQ 🎤)
 - In "Output Options":
 - "Output JSON report": Yes

Alignment with Map with BWA-MEM

BWA-MEM is a widely used sequence aligner for short-read sequencing datasets such as those we are analysing in this tutorial.

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Hands-on: Align sequencing reads to reference genome

- 1. Map with BWA-MEM & with the following parameters:
 - "Will you select a reference genome from your history or use a built-in index?": Use a genome from history and build index
 - "Use the following dataset as the reference sequence": output (Input dataset)
 - "Single or Paired-end reads": Paired Collection
 - [] "Select a paired collection": output_paired_coll (output of fastp 🎤)
 - o "Set read groups information?": Do not set
 - "Select analysis mode": 1.Simple Illumina mode

Remove duplicates with **MarkDuplicates**

MarkDuplicates Fremoves duplicate sequences originating from library preparation artifacts and sequencing artifacts. It is important to remove these artefactual sequences to avoid artificial overrepresentation of single molecule.

Hands-on: Remove PCR duplicates

- 1. **MarkDuplicates** \checkmark with the following parameters:
 - "Select SAM/BAM dataset or dataset collection": bam_output (output of Map with BWA-MEM ﴾)
 - "If true do not write duplicates to the output file instead of writing them with appropriate flags set": Yes

Generate alignment statistics with **Samtools stats**

After the duplicate marking step above we can generate statistic about the alignment we have generated.

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Hands-on: Generate alignment statistics

- 1. **Samtools stats** \(\mathcal{L} \) with the following parameters:
 - [] "BAM file": outFile (output of MarkDuplicates ﴾)
 - "Set coverage distribution": No
 - \circ "Output": One single summary file
 - o "Filter by SAM flags": Do not filter
 - "Use a reference sequence": No
 - o "Filter by regions": No

Realign reads with lofreq viterbi

Realign reads \nearrow corrects misalignments around insertions and deletions. This is required in order to accurately detect variants.

Hands-on: Realign reads around indels

- 1. **Realign reads** with lofreq \mathcal{F} with the following parameters:
 - ① "Reads to realign": outFile (output of MarkDuplicates ﴾)
 - o "Choose the source for the reference genome": History
 - **Difference**: output (Input dataset)
 - o In "Advanced options":
 - "How to handle base qualities of 2?": Keep unchanged

Add indel qualities with lofreq **Insert** indel qualities

This step adds indel qualities into our alignment file. This is necessary in order to call variants using **Call variants** with lofreq

Hands-on: Add indel qualities

- 1. Insert indel qualities with lofreq ${\cal F}$ with the following parameters:
 - \circ \square "Reads": realigned (output of Realign reads \mathcal{L})
 - "Indel calculation approach": Dindel
 - "Choose the source for the reference genome": History
 - [] "Reference": output (Input dataset)

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Call Variants using lofreq Call variants

We are now ready to call variants.

```
Hands-on: Call variants
```

- 1. **Call variants** with lofreq **\(\mu \)** with the following parameters:
 - ☐ "Input reads in BAM format": output (output of Insert indel qualities ﴾)
 - "Choose the source for the reference genome": History
 - **D "Reference": output (Input dataset)
 - o "Call variants across": Whole reference
 - o "Types of variants to call": SNVs and indels
 - o "Variant calling parameters": Configure settings
 - In "Coverage":
 - "Minimal coverage": 50
 - In "Base-calling quality":
 - "Minimum baseQ": 30
 - "Minimum baseQ for alternate bases": 30
 - In "Mapping quality":
 - "Minimum mapping quality": 20
 - o "Variant filter parameters": Preset filtering on QUAL score
 - + coverage + strand bias (lofreq call default)

The output of this step is a collection of VCF files that can be visualized in a genome browser.

Annotate variant effects with **SnpEff** eff:

We will now annotate the variants we called in the previous step with the effect they have on the SARS-CoV-2 genome.

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Hands-on: Annotate variant effects

The output of this step is a VCF file with added variant effects.

Create table of variants using **SnpSift Extract Fields**

We will now select various effects from the VCF and create a tabular file that is easier to understand for humans.

Hands-on: Create table of variants

```
1. SnpSift Extract Fields  with the following parameters:

○ " "Variant input file in VCF format": snpeff_output (output of SnpEff eff: )

○ "Fields to extract": CHROM POS REF ALT QUAL DP AF SB DP4

EFF[*].IMPACT EFF[*].FUNCLASS EFF[*].EFFECT

EFF[*].GENE EFF[*].CODON

○ "multiple field separator": ,

○ "empty field text": .
```

We can inspect the output files and see check if Variants in this file are also described in an observable notebook that shows the geographic distribution of SARS-CoV-2 variant sequences

Interesting variants include the C to T variant at position 14408 (14408C/T) in SRR11772204, 28144T/C in SRR11597145 and 25563G/T in SRR11667145.

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Summarize data with MultiQC

We will now summarize our analysis with MultiQC, which generates a beautiful report for our data.

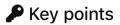
Hands-on: Summarize data

```
1. MultiQC with the following parameters:
     o In "Results":
           ■ # "Insert Results"
                 ■ "Which tool was used generate logs?": fastp
                       ■ **Dutput of fastp": report_json (output of
                        fastp 🔑)
           ■ # "Insert Results"
                 ■ "Which tool was used generate logs?": Samtools
                       ■ In "Samtools output":
                            ■ # "Insert Samtools output"
                                  ■ "Type of Samtools output?":
                                     stats
                                        ■ **D "Samtools stats output":
                                          output (output of
                                          Samtools stats \( \mathcal{L} \)
           ■ # "Insert Results"
                 "Which tool was used generate logs?": Picard
                       ■ In "Picard output":
                            ■ # "Insert Picard output"
                                  ■ "Type of Picard output?":
                                     Markdups
                                  ■ [] "Picard output": metrics_file
                                    (output of MarkDuplicates )
           ■ # "Insert Results"
                 "Which tool was used generate logs?": SnpEff
                       ■ **Output of SnpEff": csvFile (output of
                        SnpEff eff: ≯)
```

Conclusion

Congratulations, you now know how to import sequence data from the SRA and how to run an example analysis on these datasets.

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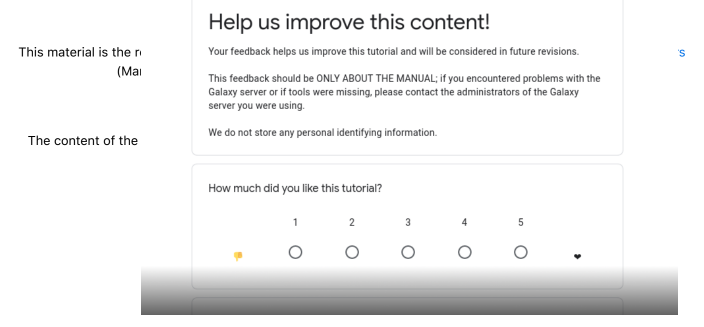
• Sequence data in the SRA can be directly imported into Galaxy

Useful literature

Further information, including links to documentation and original publications, regarding the tools, analysis techniques and the interpretation of results described in this tutorial can be found here.

Feedback

Did you use this material as an instructor? Feel free to give us feedback on how Congratulations on successfully completing this tutorial!



Citing this Tutorial

- Marius van den Beek, Dave Clements, Daniel Blankenberg, Anton Nekrutenko, 2021 From NCBI's Sequence Read Archive (SRA) to Galaxy: SARS-CoV-2 variant analysis (Galaxy Training Materials). /training-material/topics/variant-analysis/tutorials/sars-cov-2/tutorial.html Online; accessed Thu Feb 18 2021
- 2. Batut et al., 2018 Community-Driven Data Analysis Tra OPEN CHAT

Cell Systems 10.1016/j.cels.2018.05.012

BibTeX				

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