

Introduction to RNA-seq using High-Performance Computing (HPC)

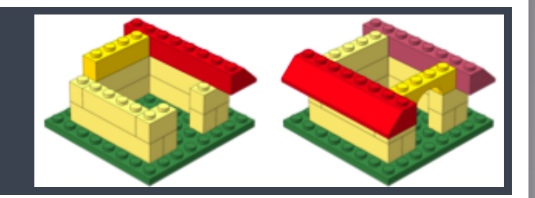
Harvard Chan Bioinformatics Core

in collaboration with

HMS Research Computing

https://tinyurl.com/hbc-rnaseq

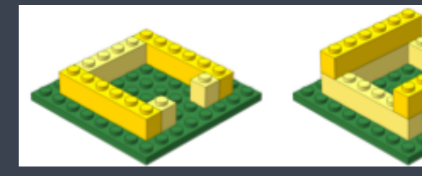
Learning Objectives

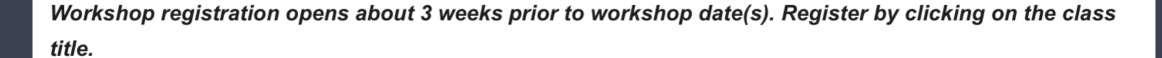


- ✓ Describe best practices for designing a bulk RNA-seq experiment
- ✓ Describe steps in an RNA-seq analysis workflow (from sequence data to expression quantification).
- ✓ Implement shell scripts on a high-performance compute cluster to perform the above steps.

We won't be covering how to perform differential gene expression (DGE) analysis on count data in this workshop.

DGE and scRNA-seq

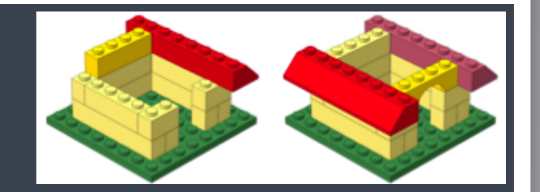




| Topic | Category | Date | Duration | Prerequisites |
|---|----------|---------------------------------|---------------------|---------------|
| Introduction to R | Basic | August 9th, 12th, 16th, 19th | Four 2h sessions | None |
| Bulk RNA-seq (Part 1 - FASTQ to counts) | Advanced | August 26th, 30th, Sept 2nd | Three 2.5h sessions | Shell |
| Bulk RNA-seq (Part II - Differential gene expression) | Advanced | Sept 9th, 13th, 16th, 20th | Four 2h session | R |
| Introduction to scRNA-seq | Advanced | Sept 27th, 30th, Oct 4th | Three 2.5h sessions | R |

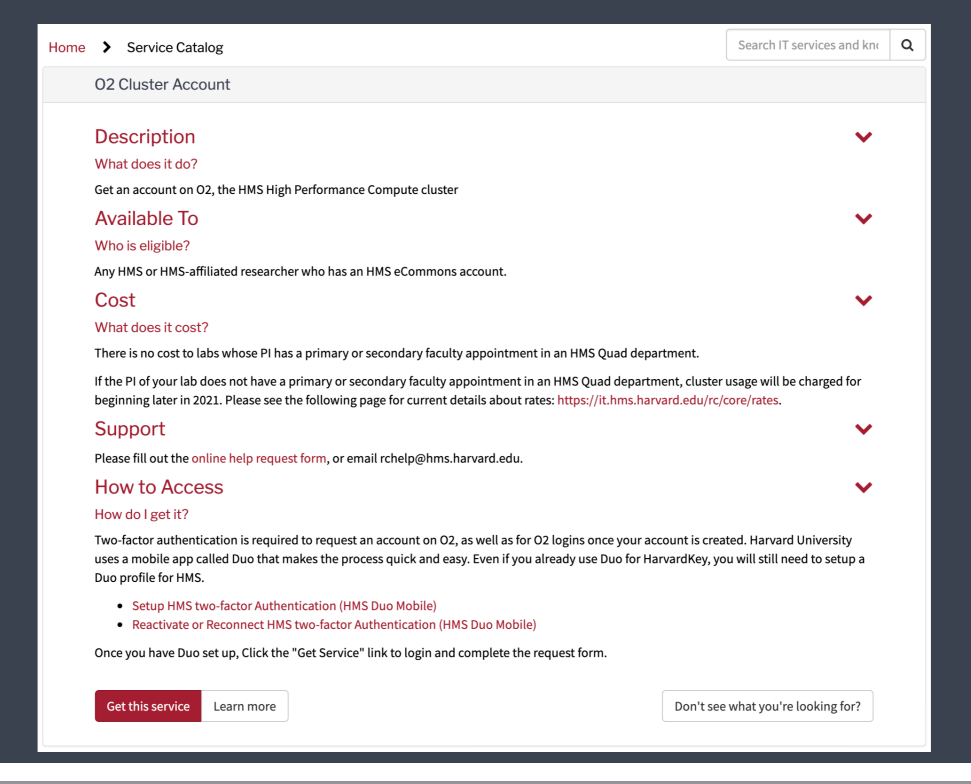
https://bioinformatics.sph.harvard.edu/upcoming-workshops

Survey



https://tinyurl.com/hbc-rnaseq-exit

Get an O2 account!



Interested in additional training?

https://hbctraining.github.io/Training-modules/

Short workshops: Current Topics in Bioinformatics

These workshops are free and open to all researchers at Harvard University and affiliated institutions.

- Workshops on bioinformatics methods & related skills.
- · Once a month for 3 hours
- · Hands-on workshops be prepared with your MAC or Windows computer
- Free and open to everyone at Harvard University and its affiliates
- · Will meet the first Wednesday of the month (with one exception) online via Zoom
- Sign up at the links below to receive the workshop Zoom link

Interested in additional training?

https://hbctraining.github.io/Training-modules/

Summer-Fall 2022 Schedule (1pm - 4pm):

| Topic and Link(s) to lessons | Prerequisites | Date | Registration |
|--------------------------------------|---|-----------|--------------|
| Basics of Python | None | 8/10/2022 | Closed |
| R practice exercises | Beginner R or Online R course - Harvard Catalyst | 9/7/2022 | Sign up! |
| Generating reports with Rmarkdown | Beginner R or Online R course - Harvard Catalyst | 10/5/2022 | Sign up! |
| Introduction to Shell | None | 11/2/2022 | Sign up! |
| Version control using Git/Github | Introduction to Shell | 12/7/2022 | Sign up! |

Interested in additional training?

All workshop materials available at:

https://hbctraining.github.io/main

Upcoming relevant courses from our partners at Countway Library:

- 1. Managing Research Data Efficiently (September 14th)
- 2. <u>Getting the Most out of Your Data: Standards to Find, Interpret & Share Data</u> (September 28th)
- 3. Achieving FAIR Data: Selecting a Repository for Your Data (October 26th)

Thanks!

- Kathleen Chappell and Andy Bergman from HMS-RC
- Data Carpentry

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