Welcome to the November 2021 Virtual CCDL scRNA-Seq Training Workshop!

Nov 1 – Nov 5, 2021 Childhood Cancer Data Lab https://alexslemonade.github.io/2021-november-training





Tell us about you!

- What's your name?
- What are you studying?
- What's a recent piece of media (book, movie, TV show, etc.)
 you enjoyed?



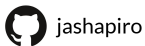
JOSH Joshua Shapiro

Data Scientist @ the CCDL

PhD Ecology & Evolution, *UChicago*Postdoc Integrative Genomics, *Princeton*

Research interests:

- Evolutionary genomics
- Single cell workflows





CHANTE
Chante Bethell

Biological Data Analyst @ the CCDL

Bachelor's in Bioinformatics from Rowan University

Research interests:

• Functional motifs in the proteome





Ally Hawkins

Data Scientist @ the CCDL

PhD Cancer Biology *University of Michigan* Postdoc Computational Biology *Cornell*

Research interests:

- Single cell data analysis
- Origins of pediatric solid tumors





JACLYN

Jaclyn Taroni

Director @ the CCDL

PhD Genetics *Dartmouth*Postdoc Computational Biology *UPenn*

Research interests:

- Transcriptomics in rare, complex diseases
- Unsupervised pattern extraction



Other staff you may see



JEN Jen O'Malley

Scientific Community Manager @ CCDL

- Maintains a supportive environment for pediatric cancer researchers
- Manages Data Lab communications



KURT Kurt Wheeler

Data Engineer @ CCDL

- Builds scalable systems
- Manages servers



DAVID David Mejia

Developer @ CCDL

- Make websites
- Debugs

Tell us about you!

- What's your name?
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Code of Conduct

Be kind, have fun

We value the involvement of everyone in the community. We are committed to creating a friendly and respectful place for learning, teaching, and contributing.

- Use welcoming and inclusive language
- Be respectful of different viewpoints and experiences
- Gracefully accept constructive criticism
- Focus on what is best for the community
- Show courtesy and respect towards other community members

Read the full Code of Conduct here:

https://alexslemonade.github.io/2021-november-training/code-of-conduct.html

If you at any time feel harassed or treated inappropriately, please contact ccdl@alexslemonade.org.

What you will learn (and what you won't)

What you will learn

We will introduce you to the R programming language, R Notebooks, and some reproducible research practices.

We cover pipelines for the quality control, pre-processing, and initial analysis of single-cell RNA-seq data almost entirely through hands-on exercises.

We generally elect to go broad and not deep.

Our overarching goals: To prepare you to perform "frontline" analyses of your own data, to get you more comfortable reading documentation/learning new methods on your own, and to give you tools to collaborate more effectively with analysts when needed

What you won't learn

We don't address experimental design (e.g., how many replicates you need).

We won't compare tools (e.g., Bioconductor vs. Seurat for single cell analysis).

We won't cover every feature (or assumption) of the tools we do present.

You may not be able to perform every analysis you need for your own work, particularly for complex experimental designs. For example, in the context of single-cell analysis, we do not cover integrating data from multiple samples.

We present analysis as a series of *linear steps*. In practice, it's **not**. It's important to consult analysis experts when you need to and to keep track of and report what you've done.

How do we pick what we teach?

We want methods to be or to have:

- Useful for a wide range of experimental designs, sample sizes
- Easy to use, well-documented, and consistently updated
- Solid tutorials, a sizeable user base, and responsive authors/maintainers

We have a preference for methods that integrate easily into a single workflow that can be run on a laptop (and our own personal biases as scientists).

Schedule

Monday	Wednesday	Friday
Workshop Intro Intro to R RStudio Server	Single-cell RNA-seq Dimensionality reduction Clustering Marker identification	Consultations Your own data Exercise notebooks
Base R ggplot2 & tidyverse Consultations Exercise notebooks	Consultations Exercise notebooks Your own data	<u>Presentations</u>
Tuesday	Thursday	
Single-cell RNA-seq Quantification & QC Filtering & Normalization	Pathway Analysis Overrepresentation GSEA	
Consultations Exercise notebooks Your own data	Consultations Exercise notebooks Your own data	

 $Full schedule: {\tt https://alexslemonade.github.io/2021-november-training/workshop/SCHEDULE.html}$

Daily Schedule Outline

Instruction

Full group Lectures

Breakout

Small groups
Start exercise notebooks

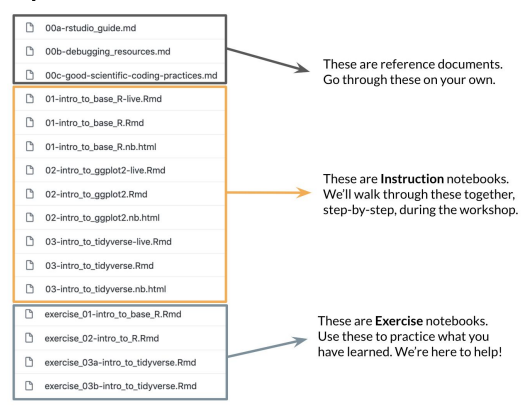
Consultation Period

Exercise notebooks Your own data

- Introduce concepts and background
- Demonstrate usage
- Answer general questions
- Split up into Zoom breakout rooms
- Ask questions of instructors and other participants

- Practice what you have learned
- Work on exercises individually or in groups
- Work with your own data

Module Layout



Module cheatsheets cover key functions

https://github.com/AlexsLemonade/training-modules/tree/2021-november/module-cheatsheets

dplyr

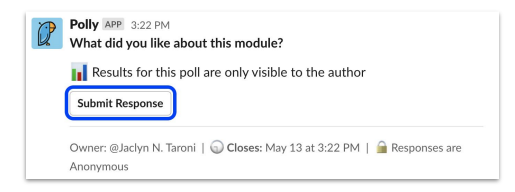
Read the dplyr package documentation here.

A vignette on the usage of the dplyr package can be found here.

Library/Package	Piece of code	What it's called	What it does
dplyr	<u>\$>\$</u>	Pipe operator	Funnels a data.frame through tidyverse operations
dplyr	filter()	Filter	Returns a subset of rows matching the conditions of the specified logical argument
dplyr	arrange()	Arrange	Reorders rows in ascending order. arrange(desc()) would reorder rows in descending order.
dplyr	select()	Select	Selects columns that match the specified argument
dplyr	mutate()	Mutate	Adds a new column that is a function of existing columns
dplyr	summarise()	Summarise	Summarises multiple values in an object into a single value. This function can be used with other functions to retrieve a single output value for the grouped values. summarize and summarise are synonyms in this package.
dplyr	rename()	Rename	Renames designated columns while keeping all variables of the data.frame
dplyr	group_by()	Group By	Groups data into rows that contain the same specified value(s)
dplyr	inner_join()	Inner Join	Joins data from two data frames, retaining only the rows that are in both datasets.

We want your feedback!

At the end of each module, we will post a few questions in the Slack channel.



- The most difficult or confusing point of the module ("muddiest point")
 We will post additional material answering your questions the next day
 Responses to this question will appear in the channel anonymously
- What did you like about the module?
- How we can improve the module?
 These responses will be collected anonymously (and not posted)

Friday

Your own projects Exercise notebooks

Spend Friday working with your own data, getting assistance as needed from CCDL staff and each other.

Presentations

Present what you worked on during the consultation times to the group!

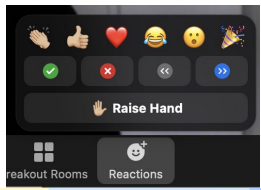
Virtual Training Procedures



General Zoom Etiquette

- Keep your microphone muted
- Type questions in the Chat window (directed to the host)

- To: Jaclyn Taroni ▼ (Privately)
 Type message here...
- Click on the "Chat" button at the bottom of your window to open the chat.
- Use the Zoom reactions to tell us how you are doing!

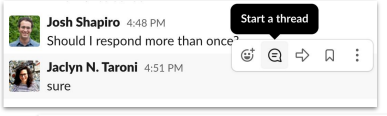


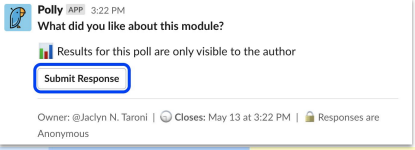


Using Slack

- Use the #2021-november-training Slack channel
- Post public questions, get help with errors and debugging, make comments, and help your fellow participants!
 - Use threads to keep related content together

Watch out for feedback polls!





Communication during instruction



- I have an urgent question that needs an answer before moving on:
 - Raise Hand or Chat with the room host
- I'm stuck with an error and can't proceed with the hands-on exercise
 - *Chat* with meeting host: Request 1:1 and you will be placed in a breakout room with a CCDL staff member



- I have an general question that does not need an answer right away.
 - **Post** in #2021-november-training
- I'm having trouble logging in to RStudio Server
 Direct Message a CCDL staff member (not the current host or instructor)

Trouble logging into Zoom and Slack? *Email* training@ccdatalab.org

Communication at other times (consultation periods)



- I have questions about previous instruction or exercise notebooks
 - **Post** in #2021-november-training
 - If you need to share your screen, we will set up a 1:1 or group Zoom call
- I would like to be paired up with other participants
 - **Post** in #2021-november-training; we can set you up in a Zoom breakout room

- I have a question that is highly specific to my data
 - **Direct Message** a CCDL staff member
- I'm having trouble logging in to RStudio Server
 - **Direct Message** a CCDL staff member

Trouble logging into Zoom and Slack? *Email* training@ccdatalab.org