

Introduction to R

Harvard Chan Bioinformatics Core

<https://tinyurl.com/hbc-r-online-nanocourse>

Sponsored by DF/HCC, HSCI, HMS Foundry



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Director



Victor Barrera



Amelie Jule



Zhu Zhuo



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Meeta Mistry



Jihe Liu



Will Gammerdinger



Maria Simoneau



James Billingsley



Sergey Naumenko



Peter Kraft
Faculty Advisor

Consulting

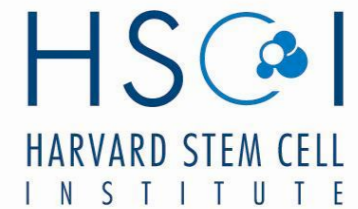
- Experimental design help
- RNA-seq analysis: bulk, single cell, small RNA
- ChIP-seq and ATAC-seq analysis
- Genome-wide methylation
- WGS, resequencing, exome-seq and CNV studies
- QC & analysis of gene expression arrays
- Functional enrichment analysis
- Grant support

<http://bioinformatics.sph.harvard.edu/>



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SCHOOL OF PUBLIC HEALTH

NIEHS



Training

We have divided our short workshops into 2 categories:

1. Basic Data Skills - No prior programming knowledge needed (no prerequisites)
2. Advanced Topics: Analysis of high-throughput sequencing (NGS) data - Certain “Basic” workshops required as prerequisites.

Any participants wanting to take an advanced workshop will have to have taken the appropriate basic workshop(s) within the past 6 months.

<http://bioinformatics.sph.harvard.edu/training/>

<https://hbctraining.github.io/main/>



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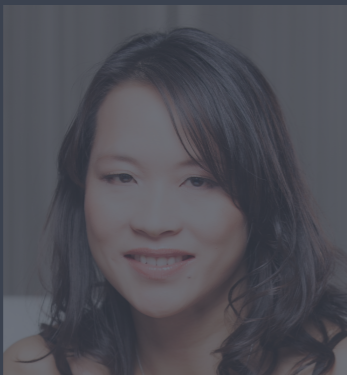
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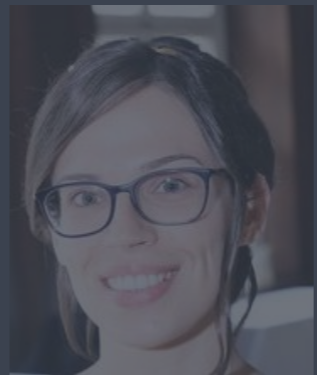
Introductions!



Shannan Ho Sui
Director



Victor Barrera



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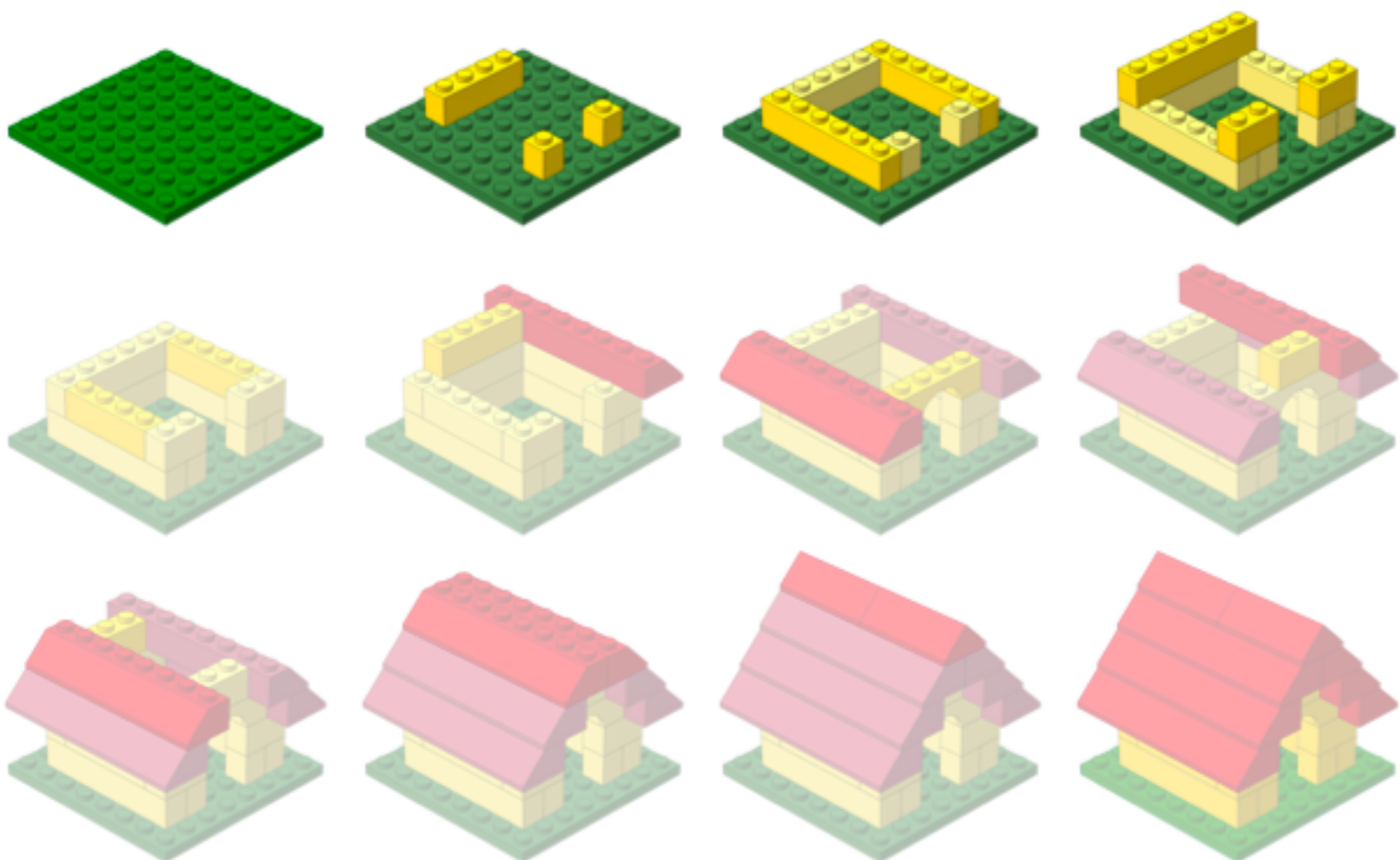


Sergey Naumenko



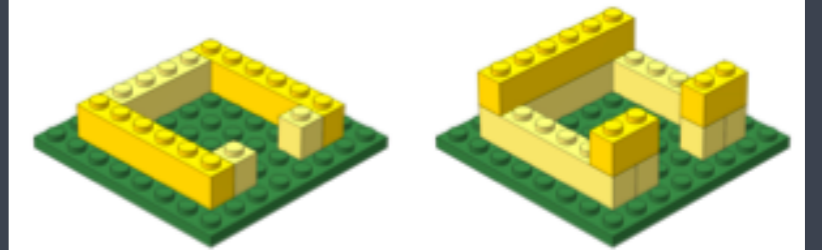
Peter Kraft
Faculty Advisor

Nanocourse Scope...



Learning R

Nanocourse Scope



- ✓ Comfortably use RStudio (a graphical interface for R)
- ✓ Fluently interact with R using RStudio
- ✓ Become familiar with R syntax
- ✓ Understand data structures in R
- ✓ Inspect and manipulate data structures
- ✓ Install packages and use functions in R

CRAN

(Comprehensive R Archive Network)



Available CRAN Packages By Name

[A](#) [B](#) [C](#) [D](#) [E](#) [F](#) [G](#) [H](#) [I](#) [J](#) [K](#) [L](#) [M](#) [N](#) [O](#) [P](#) [Q](#) [R](#) [S](#) [T](#) [U](#) [V](#) [W](#) [X](#) [Y](#) [Z](#)

A3	Accurate, Adaptable, and Accessible Error Metrics for Predictive Models
abbyyR	Access to Abbyy Optical Character Recognition (OCR) API
abc	Tools for Approximate Bayesian Computation (ABC)
ABCanalysis	Computed ABC Analysis
abc.data	Data Only: Tools for Approximate Bayesian Computation (ABC)
abcdeFBA	ABCDE_FBA: A-Biologist-Can-Do-Everything of Flux Balance Analysis with this package
ABCOptim	Implementation of Artificial Bee Colony (ABC) Optimization
ABCp2	Approximate Bayesian Computational Model for Estimating P2
abcrf	Approximate Bayesian Computation via Random Forests

*CRAN
Mirrors
What's new?
Task Views
Search

About R
R Homepage
The R Journal*

- The main repository for R packages
- Easy to install

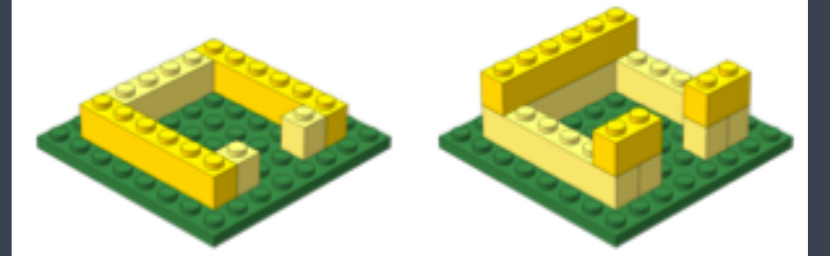
<https://cran.r-project.org/>



- An alternative package repository; “..provides tools for the analysis and comprehension of *high-throughput genomic data*.”
- Includes (but is not limited to) tools for:
 - performing statistical analysis
 - accessing public datasets
- Open source and open development
- Free

www.bioconductor.org

Nanocourse Scope



- Comfortably use RStudio (a graphical interface for R)
 - Fluently interact with R using RStudio
 - Become familiar with R syntax
 - Understand data structures in R
 - Inspect and manipulate data structures
 - Install packages and use functions in R
- ✓ Visualize data using *ggplot2*
- ✓ Utilize pipes, tibbles and functions from the Tidyverse package suite

Logistics

Course webpage

<https://tinyurl.com/hbc-r-online-nanocourse>

Course schedule online

Workshop Schedule

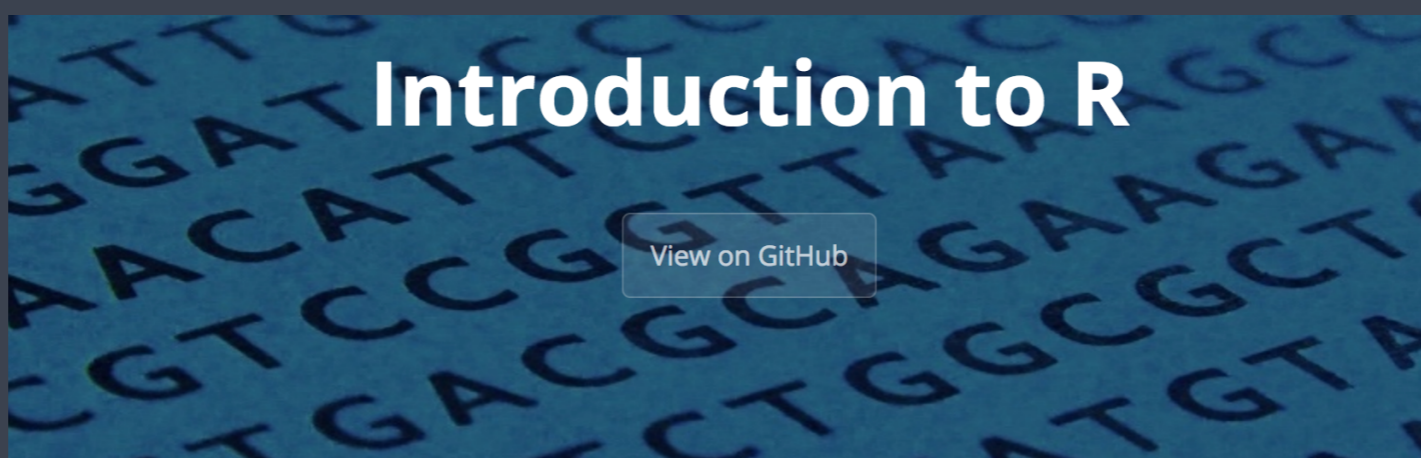
Day 1

Time	Topic	Instructor
10:00 - 10:30	Workshop Introduction	Jihe
10:30 - 11:45	Introduction to R and RStudio	Radhika
11:45 - 12:00	Overview of self-learning materials and homework submission	Mary

Before the next class:

1. Please **study the contents** and **work through all the code** within the following lessons:
 - o [R Syntax and Data Structure](#)
 - o [Functions and Arguments](#)
 - o [Reading in and inspecting data](#)
2. **Complete the exercises:**
 - o Each lesson above contain exercises; please go through each of them.
 - o **Copy over** your code from the exercises into a text file.
 - o **Upload the saved text file** to [Dropbox](#) the **day before the next class**.

Course materials online



Introduction to R

[View on GitHub](#)

Approximate time: 70 min

Learning Objectives

- Employ variables in R.
- Describe the various data types used in R.
- Construct data structures to store data.

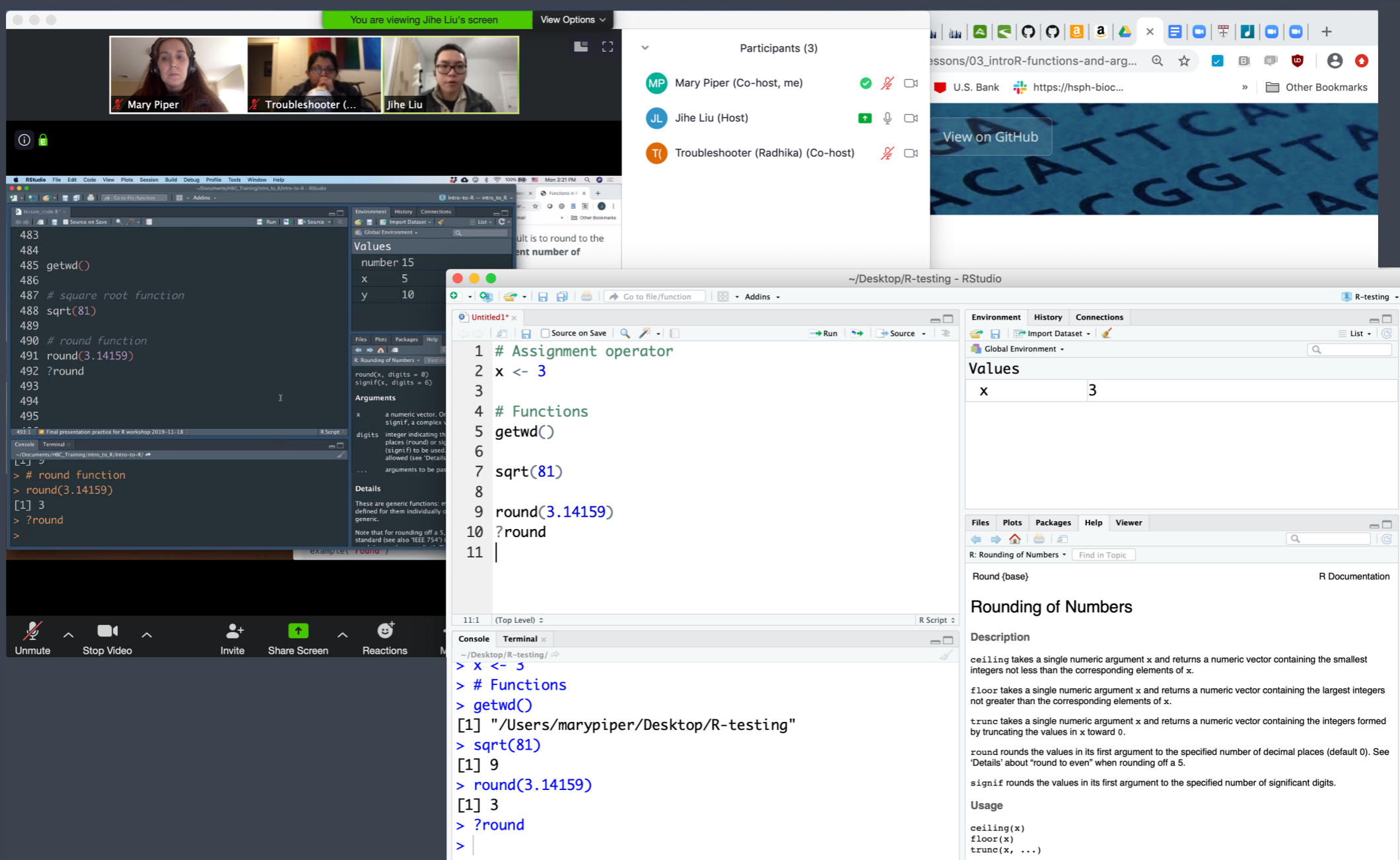
The R syntax

Now that we know how to talk with R via the script editor or the console, we want to use R for something more than adding numbers. To do this, we need to know more about the R syntax.

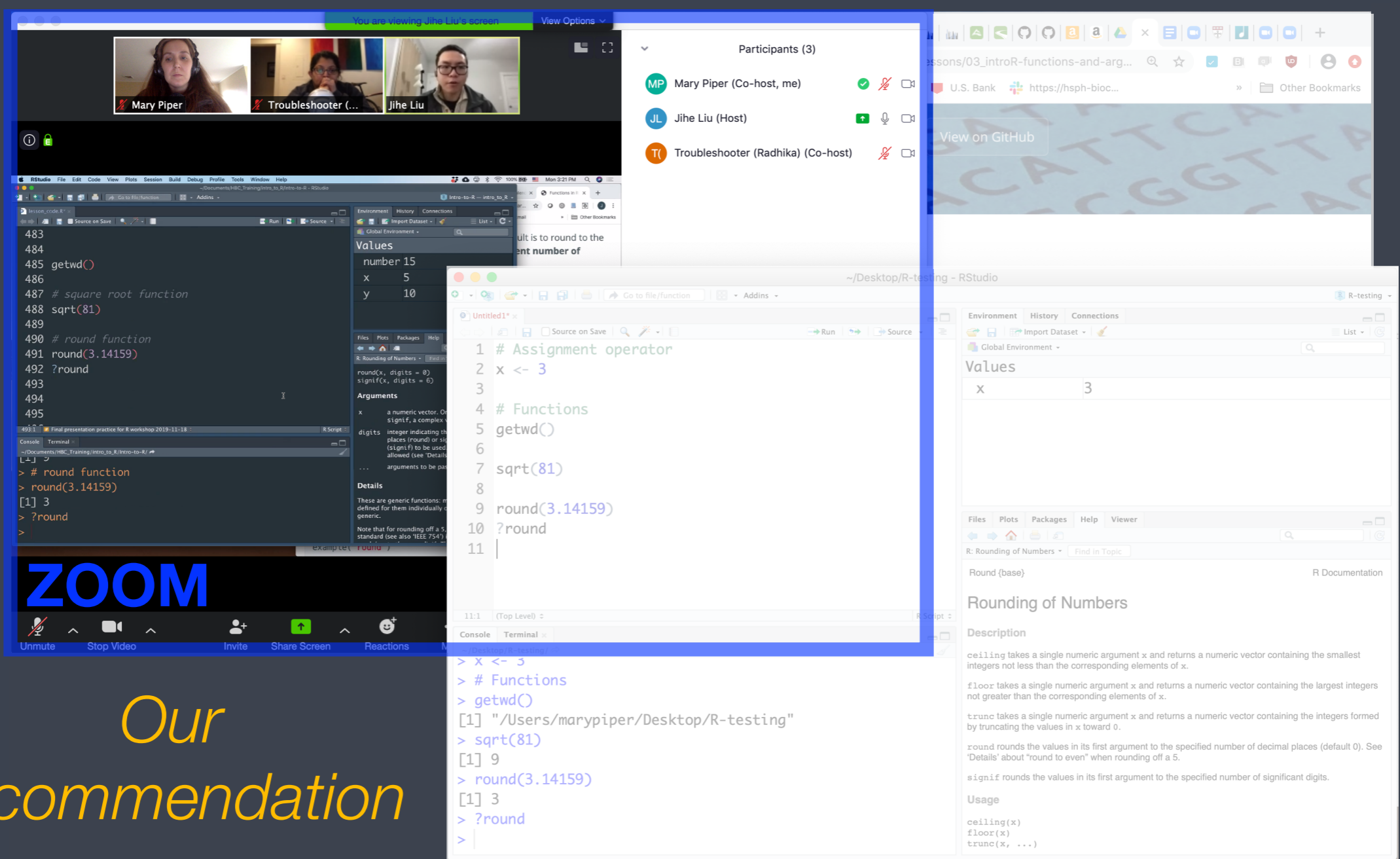
Below is an example script highlighting the many different “parts of speech” for R (syntax):

- the **comments** `#` and how they are used to document function and its content
- **variables and functions**
- the **assignment operator** `<-`

Single screen & 3 windows?

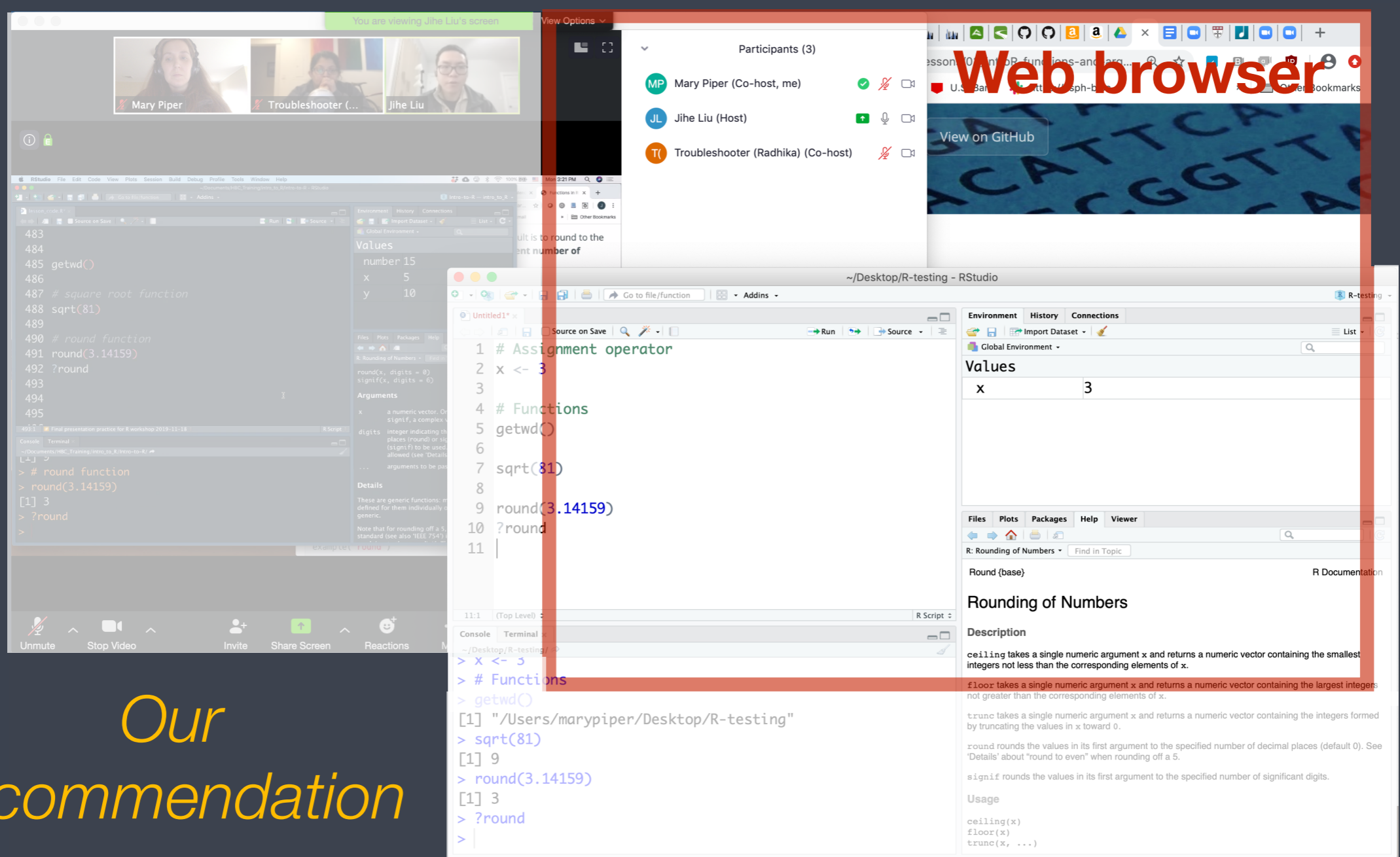


Single screen & 3 windows?



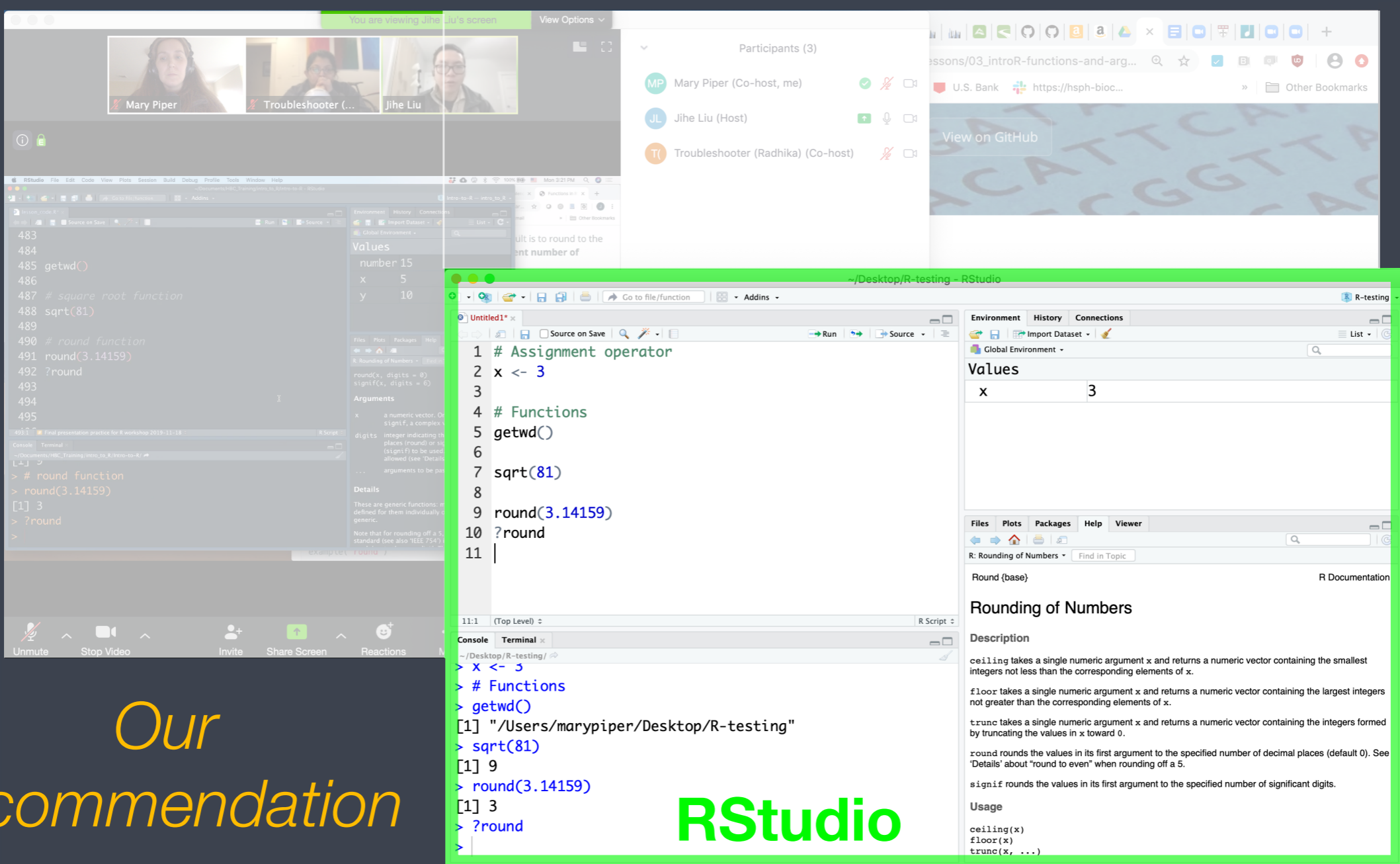
*Our
recommendation*

Single screen & 3 windows?

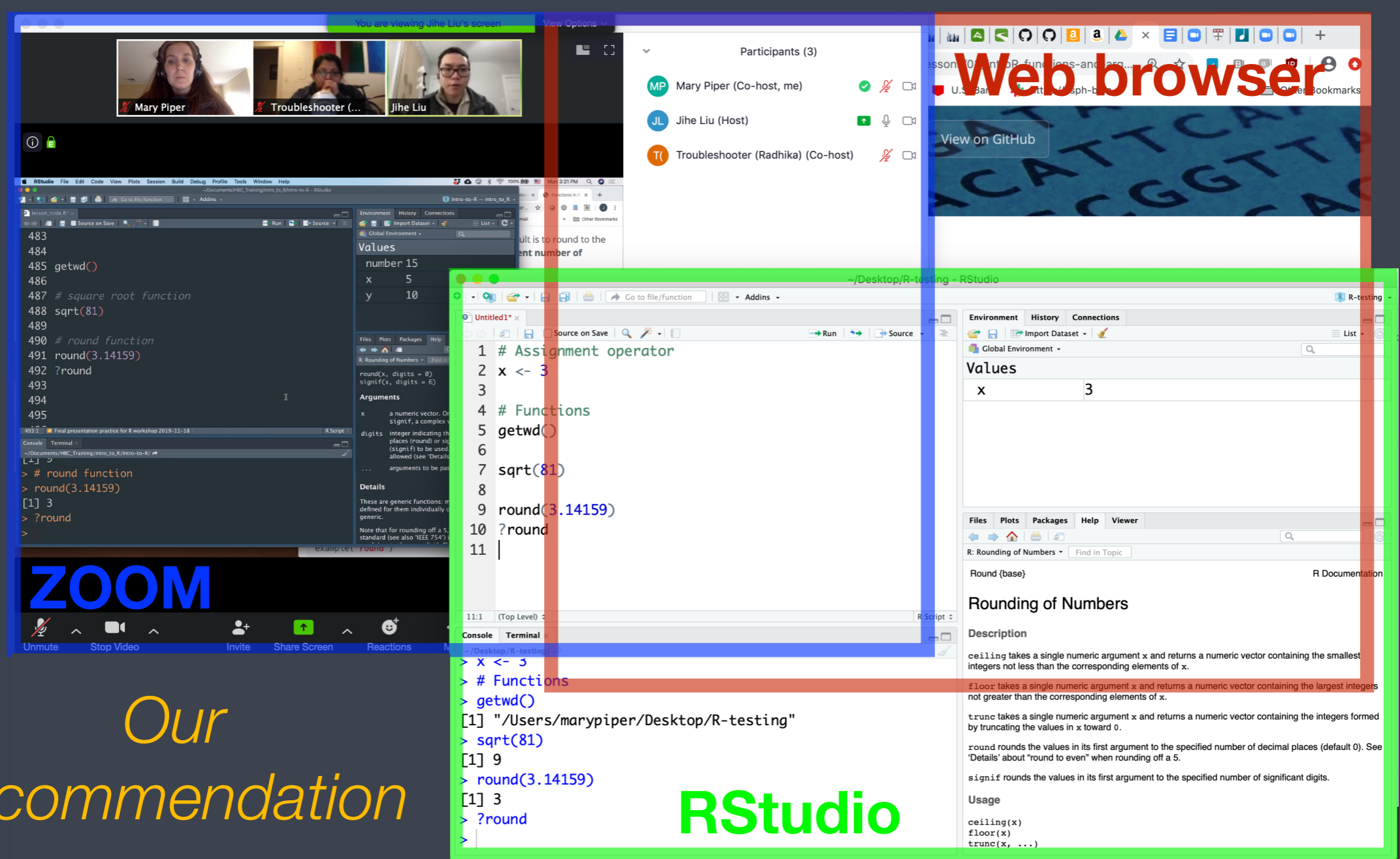


*Our
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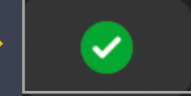
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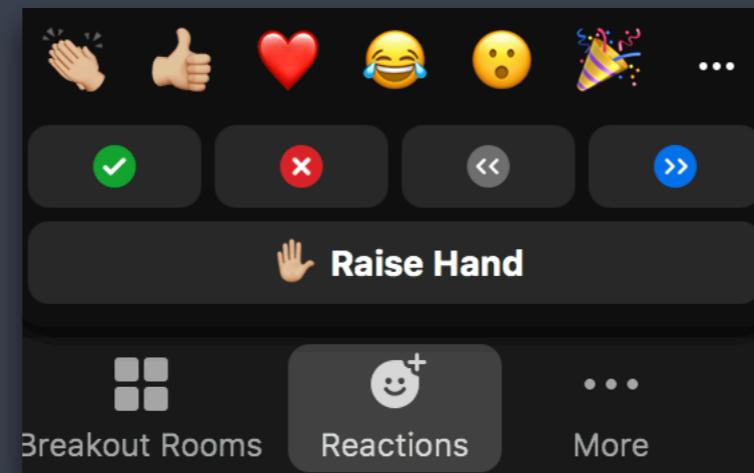


Odds and Ends

- ❖ Quit/minimize all applications that are not required for class
- ❖ Captioning is available upon request

Odds and Ends (1/2)

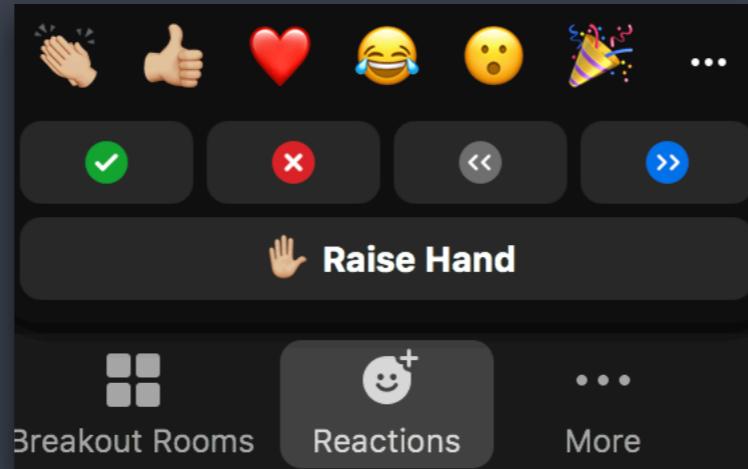
- ❖ Quit/minimize all applications that are not required for class
- ❖ Captioning is available upon request
- ❖ Are you all set?
 - ▶  = "agree", "I'm all set" (equivalent to a **green post-it**)
 - ▶  = "disagree", "I need help" (equivalent to a **red post-it**)



Odds and Ends (2/2)

- ❖ Questions for the presenter?

- Post the question in the Chat window OR
-  when the presenter asks for questions
- Let the Moderator know



Credit Considerations

- ❖ To receive credit for this course:
 - ❖ Attend all sessions in their entirety
 - ❖ Complete all self-learning exercises

Contact us!

HBC training team: hbctraining@hsph.harvard.edu

HBC consulting: bioinformatics@hsph.harvard.edu

Twitter

[@bioinfocore](https://twitter.com/bioinfocore)