

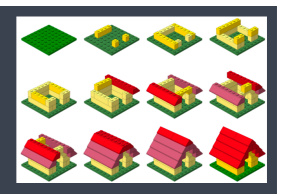


Introduction to Differential Gene Expression Analysis

Harvard Chan Bioinformatics Core

https://tinyurl.com/hbc-intro-to-dge

Workshop Scope



- Understand the considerations for performing statistical analysis on RNA-seq data
- Start with gene counts (after alignment and counting)
- Perform QC on count data
- Use DESeq2 to perform differential expression analysis on the count data and obtain a list of significantly different genes
- Visualize results of the analysis
- Perform functional analysis on the lists of differentially expressed genes

Resources

- All workshop materials
- <u>2-3 hour workshops</u>
- DESeq2 vignette
- More about color palettes in R
- Multiple plots in a single figure using gridExtra

Exit survey

http://tinyurl.com/intro-to-dge-exit

Bioinformatics Community Networking Breakfast!

- Free and open to the LMA community
- Food and seats are first-come-first-served
- Last Wednesday of every month

Date: October 30th, 2019

Time: 9:00 to 10:30 am

Venue: Modell 100A (Armenise)

Fred S. Rosen Lecture Hall



Contact us!

HBC webpage: http://bioinformatics.sph.harvard.edu

Training email: hbctraining@hsph.harvard.edu

Consulting email: bioinformatics@hsph.harvard.edu

Twitter

@bioinfocore