

Introduction to Bioinformatics and Computational Biology

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Chapter 1

Course information

This is the course material for STAT115/215 BIO/BST282 at Harvard University.

All the YouTube videos in this course are organized under the 2021 STAT115 playlist.

1.1 Contributors

Xiaole Shirley Liu Harvard University and Dana-Farber Cancer Institute

Joshua Starmer StatQuest

Ting Wang Washington University

Feng Yue Northwestern University

Ming Tang Dana-Farber Cancer Institute

Martin Hemberg Wellcome Sanger Institute

Gad Getz Harvard University and Broad Institute

Yang Liu

Bo Yuan

Jack Kang

Scarlett Qian

Jiazhen Rong

Phillip Nicol

Maartin De Vries

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Chapter 2

Introduction

2.1 Brief history of bioinformatics

2.1.1 Protein structure wave

2.1.2 Gene expression wave

2.1.3 Genome sequencing wave

2.1.4 Big data challenge from sequencing

2.2 Should I take this course?

2.2.1 Bioinformatics vs computational biology

2.2.2 Is this class for me?

2.3 Course information

2.3.1 Logistics

2.3.2 X Shirley Liu lab introduction

2.4 Lab 1

2.4.1 Introduction**2.4.2 Introduction to R****2.4.3 Introduction to Bash****2.4.4 Getting started with Cannon**

Chapter 3

High throughput sequencing

3.1 Three generations of sequencing technologies

First generation sequencing is Sanger sequencing. It is the technology that was used to obtain the first human genome sequence.

Second generation sequencing is also called next generation sequencing (NGS) and is the start of high throughput sequencing. It is what scientists use most often nowadays, and Illumina is the market leader. Most of the rest of this course will cover data analysis using second generation sequencing.

Third generation sequencing is single-molecule sequencing. There are many new technologies still under active development, although none has reached market penetration.

3.2 FASTQ and FASTQC

NGS generates FASTQ files. FASTQC is an computational approach to evaluate the quality of your NGS data.

3.3 Early sequence alignment (1 with 1)

In the early days (1970s), scientists were not worried about having to align too many sequences. They wanted to find the best alignment between two

sequences. Many bioinformatics courses start with learning these, although it is not the main focus of our course. We included two videos in case you are interested.

The Needleman-Wunsch algorithm is the earliest algorithm to find the alignment between two sequences and score their similarity.

When two sequences are long, and only a portion of them can align well with each other, the Smith-Waterman algorithm can find the best local sequence alignment. It is still considered the best alignment approach, although it is slow.

3.4 Sequence search algorithms (1 with many)

With more and more sequences available in the public in the 1980s, scientists were interested in finding whether their newly sequenced string has been sequenced before in the public database. Therefore, the fast search algorithm BLAST was developed, using one sequence as the query to find similar sequences from a database.

3.5 Borrow-Wheeler Aligner (many with many)

With NGS, scientists need much faster search (aka mapping) algorithms in order to align the millions of sequences to the reference genome. The current best algorithm is called Borrow-Wheeler Aligner or BWA.

In order to understand BWA, we first need to introduce Borrow-Wheeler transformation and LF mapping

The basic idea of Borrow-Wheeler alignment

3.6 Alignment output

NGS raw data is in FASTQ. Alignment gives you SAM (alignment) or BAM (binary version of SAM) files which contain the sequence information in FASTQ and the mapping locations. BED file is the simplest, although there is information loss.

Chapter 4

RNA-seq Quantification

- 4.1 Introduction to RNA-seq experiment
- 4.2 RNA quality control and experimental design
- 4.3 Alignment
- 4.4 RNA-seq QC
- 4.5 RNA-seq expression index
- 4.6 RSEM and Salmon
- 4.7 RNA-seq read distribution
- 4.8 Lab 2
 - 4.8.1 STAR tutorial
 - 4.8.2 RSeQC tutorial
 - 4.8.3 RSEM/Salmon Tutorial

Chapter 5

Differential expression, FDR, GO, and GSEA

DESeq2 is a popular and accurate computational algorithm to detect differential gene expression from RNA-seq data. It includes many elegant quantitative considerations, such as:

- Normalize the gene read counts by library size and composition
- Model gene read counts with negative binomial distribution
- Use hierarchical modeling to stabilize the gene variance
- Use Benjamini-Hochberg to calculate control for false discovery rate of calling differentially expressed genes
- Filter lowly expressed genes to reduce the number of hypotheses to be tested

5.1 DESeq2 library normalization

5.2 DESeq2 variance stabilization

5.3 Multiple hypotheses testing and False Discovery Rate

5.4 DESeq2 gene filtering

5.5 Gene Ontology (GO analysis)

5.6 Gene Set Enrichment Analysis (GSEA)

5.7 DESeq2 tutorial

Chapter 6

Clustering

6.1 Heatmap and clustering quality

6.2 Hierarchical cluster

6.3 K means cluster

6.4 Pick K and consensus clustering

6.5 Batch effect removal

6.6 Lab3

6.6.1 PCA tutorial

6.6.2 Clustering tutorial

6.6.3 Combat tutorial

6.6.4 DESeq2 Tutorial

6.6.5 DAVID/GSEA Tutorial

Chapter 7

Dimension Reduction

RNA-seq samples have tens of thousands of genes, although many genes might not vary much between samples and many others have correlated gene expression. Dimension reduction techniques aim to reduce the dimension of representing each sample with tens of thousands of genes to much fewer dimensions, e.g. 2 to 100.

7.1 Principal Component Analysis: idea behind PCA.

PCA / SVD automatically outputs PC1, PC2, PC3, etc, with earlier PCs capturing the highest level of variability in the original data. Each PC is a linear combination of raw gene expression, and is orthogonal to all other PCs.

7.2 Principal Component Analysis: PCA applications.

PCA is a widely used method to project samples with high dimensions (e.g. with gene expression data) onto two dimensions for better visualization. It is an intuitive way to identify sample clusters, and identify batch effect.

7.3 Multidimensional Scaling (MDS)

MDS can use differential ways to calculate pair-wise distance, then use lower dimensions to satisfy the pair-wise distance. PCA is a special case of MDS.

7.4 Linear discriminant Analysis (LDA)

LDA is not only a dimension reduction method, but also a supervised machine learning method.

Chapter 8

Classification

8.1 Introduction

Imagine you have RNA-seq of a collection of labeled normal lung and lung cancer tissues. Given a new sample of RNA-seq from the lung with unknown diagnosis, will you be able to predict based on the existing labeled samples and the expression data whether the new sample is normal or tumor? This is a sample classification problem, and it could be solved using **unsupervised** and **supervised** learning approaches.

Unsupervised learning is basically clustering or dimension reduction. You can use hierarchical clustering, MDS, or PCA. After clustering and projection the data to lower dimensions, you examine the labels of the known samples (hopefully they cluster into separate groups by the label). Then you can assign label to the unknown sample based on its distance to the known samples.

Supervised learning considers the labels with known samples and tries to identify features that can separate the samples by the label. Cross validation is conducted to evaluate the performance of different approaches and avoid over fitting.

StatQuest has done an amazing job with machine learning with a full playlist of well organized videos. While the full playlist is worth a full course, for the purpose of the course, we will just highlight a number of widely used approaches. They include logistic regression (this is considered statistical machine learning), K nearest neighbors, random forest, and support vector machine (these are considered computer science machine learning).

8.2 Supervised learning

8.3 Cross validation

8.4 Regression

8.5 Regularization

8.5.1 Ridge regression

8.5.2 LASSO regression

8.5.2.1 LASSO tutorial in R

8.6 KNN

8.7 Decision trees

8.8 Random forest

8.9 SVM

8.10 Lab 4

8.10.1 K-Nearest Neighbors tutorial

8.10.2 Regression/Ridge/LASSO Tutorial

8.10.3 Logistic Regression Tutorial

8.10.4 Support Vector Machine Tutorial

8.10.5 Random Forest Tutorial

Chapter 9

Module I Review

9.1 Module I review

9.2 Analysis Scenario 1

9.3 Analysis Scenario 2

Chapter 10

Transcription Factor Motif Finding

10.1 Transcription regulation

10.2 Motif representation

10.3 EM

10.4 Gibbs sampler

10.5 Gibbs intuition

10.6 Motif finding in eukaryotes

10.7 Known motif database

Chapter 11

ChIP-seq, Expression Integration

11.1 Motif finding in eukaryotes, and ChIP-seq

11.2 MACS and ChIP-seq QC

11.3 Identify TF interactions from ChIP-seq motifs

11.4 TF target genes and expression integration

11.5 Lab 5

11.5.1 MACS Tutorial

11.5.2 ChIP-seq QC Tutorial

11.5.3 TF Motif Finding Tutorial

11.5.4 TF Collaborator Tutorial

Chapter 12

Epigenetics, DNA Methylation

12.1 Intro to DNA Methylation

12.2 DNA Methylation Pattern and Function

12.3 DNA Methylation in Diseases

12.4 Techniques to Measure DNA Methylation

Chapter 13

Histone Modifications , Chromatin Accessibility

13.1 Nucleosome Positioning

13.2 Introduction to Histone Modifications

13.3 Infer Transcription Factor Binding from
Histone Mark Dynamics

13.4 Using Histone Marks to Infer Gene Func-
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13.5 Introduction to DNase-seq and ATAC-seq

13.6 Infer TF from Differential Genes Using
LISA

13.7 Caution on DNase/ATAC-seq footprint
analysis

13.8 Summary of Epigenetics and Chromatin

13.9 Lab 6

13.9.1 ChIP-seq Expression Integration

13.9.2 Cistrome-GO Tutorial

13.9.3 ATAC-seq Analysis and LISA Tutorial

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Hidden Markov Model

14.1 Markov Chain

14.2 Hidden Markov Model

14.3 Hidden Markov Model Forward Procedure

14.4 Hidden Markov Model Backward Procedure

14.5 HMM Forward-Backward Algorithm

14.6 Viterbi Algorithm

14.7 Baum Welch Algorithm Intuition

14.8 HMM Bioinformatics Applications

Chapter 15

HiC

- 15.1 Introduction to Chromatin Interaction and Organization
- 15.2 Methods to Investigate 3D Genome Organization
- 15.3 Topologically Associating Domains
- 15.4 TAD Function and Loop Anchors
- 15.5 Chromatin Compartments
- 15.6 Computational Methods to Call Chromatin Loops
- 15.7 Variations of Chromatin Interaction Technologies
- 15.8 Resources for Exploring 3D Genomes

15.9 Lab 7

15.9.1 BS-seq and Bismark Tutorial

15.9.2 Tutorial on Associating DNA Methylation with Expression

15.9.3 HiC Analysis Tutorial

Chapter 16

Module II Review

16.1 Module II Review

16.2 Module II Analysis Scenarios

Chapter 17

SNP and GWAS

17.1 SNP, LP, and Association Studies

17.2 GWAS Studies and eQTL Analysis

17.3 Lab 8

17.3.1 HW4 FAQ & cooler

17.3.2 Pikachu&HiGlass

17.3.3 HMM

Chapter 18

GWAS and Epigenomics

18.1 Intro Functional Annotate GWAS

18.2 GWAS Functional Enrichment

18.3 Find Causal SNPs

18.4 Predict disease risk

Chapter 19

Single-cell RNA-seq (1)

19.1 Intro to scRNA-seq

19.2 scRNA seq techniques

19.3 scRNA seq preprocessing and QC

19.4 Cleaning up expression matrix

Chapter 20

Single-cell RNA-seq (2)

20.1 scRNA seq dimension reduction

20.2 Clustering and projections

20.3 Pseudo time and RNA velocity

20.4 Clustering by genotype and CITE seq

Chapter 21

scATAC-seq

21.1 Single-Cell ATAC-seq Technique

21.2 Single-Cell ATAC-seq Pre-Processing and QC

21.3 Single-Cell ATAC-seq Analysis

21.4 scATAC-seq Downstream Analyses and scRNA-seq Integration

21.5 Lab 9

21.5.1 MAESTRO tutorial

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Module III Review

22.1 Module III Review

Chapter 23

Cancer Genome Sequencing , Mutation analyses

23.1 Intro to TCGA

23.2 Cancer mutation characterization

23.3 Cancer mutation patterns

23.4 Tumor purity and clonality

23.5 Interpret tumor mutations

23.6 Find cancer genes

23.7 Summary and future

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Cancer Subtyping, Survival Analyses

24.1 Tumor Subtypes

24.2 Survival analysis

24.3 Oncogenes and Tumor Suppressor Mutations

24.4 Cancer Epigenetics

Chapter 25

Targeted Therapy and Precision Medicine

- 25.1 Introduction to Targeted Therapy and Precision Medicine
- 25.2 Resistance to targeted therapy
- 25.3 Model system chemical and genetic screens
- 25.4 Overcoming resistance to targeted therapy

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Cancer Immunotherapy (1)

26.1 Intro to Cancer Immunotherapy

26.2 HLA and Neoantigen Presentation

26.3 Immune Cell Infiltration in Tumors

26.4 T Cell Receptor Repertoires in Cancer Immunology

26.5 Lab 10

26.5.1 TCGA exploration

26.5.2 LIMMA on microarray data

26.5.3 Survival analysis

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Cancer Immunotherapy (2)

27.1 B Cell Receptor Repertoires in Tumors

27.2 T Cell Activation and Dysfunction

27.3 NK Cells and Macrophages in Tumor Immunity

27.4 Cancer Immunotherapy Response Biomarkers

27.5 Improving Immunotherapy Response

27.6 Lab 11

27.6.1 Cancer mutations and driver genes

27.6.2 CRISPR screen

27.6.3 Cancer immunology

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CRISPR Screens

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28.2 CRISPRa and CRISPRi

28.3 CRISPR design and outcome

28.4 CRISPR screens & DepMap

28.5 CRISPR screen analysis

28.6 CRISPR screens in drug response

28.7 CRISPR screens in immunology

28.8 Enhancer CRISPR screen

28.9 CRISPR screens + scRNA-seq

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Module IV Review and Course Review

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29.2 Course Review