Statistical Methods for High Dimensional Biology

STAT/BIOF/GSAT 540

Lecture I – course introduction
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Today's topics

- What the course is about
- Course mechanics
- Introduction to high-dimensional biology

Your instructors

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Course audience

- Researchers who want to know how to analyze large data sets from biological studies
- Genomics-focused, but information is broadly applicable
- Statistics students might find the math parts easy
- Biology students might find the biology easy
- We are counting on you to help make it work: help your peers!

Prerequisites

Officially, none. But:

- Statistics You should have already taken university level "Statistics 101". You'll get a refresher, but you should be prepared to get comfortable thinking about things like "probabilities" and "specificity".
- Biology No requirements, but you are expected to learn things like the difference between a DNA and RNA and a gene and a genome. We assume you are here because you are interested in biology and will pick it up.
- No R experience required but you must be prepared to do a lot of self-guided learning.
- You'll use your own computer to run R. If you can't install R on your computer, ask us for options.

What you can expect to learn

- Conceptual and practical knowledge you need to handle large biological data sets
 - Less about specific types of data, more about generally applicable approaches and principles
- You will be able to critically evaluate analyses in the literature
- Implementation of analyses using the R/Bioconductor computing environment

Not about:

- Formal mathematical theory underpinning the approaches
- Gory details of how to analyze any particular type of data at a low level

Topics covered

Probability foundations

Exploratory data analysis

Data QC and preprocessing

Basic statistical inference ("one gene at a time")

Large-scale inference ("genome-wide")

Count-based data (e.g. RNA-seq) analysis

DNA methylation analysis

Principal Component Analysis

Clustering

Classification

Resampling and bootstrap

Model selection and regularization

Gene sets and gene networks

Course mechanics

Course web site

http://stat540-ubc.github.io/

- Lecture notes
- Lab notes
- Assignments

Much interaction via Github (discussions, submission)

https://github.com/STAT540-UBC

Lectures

- Lectures shared among three professors
- Notes provided on web before class

Sections/Labs

- Wednesdays in room ESB 1042
- Officially from 12-1, but we will start at 11
 - 11-12: Lab available
 - 12-1: TA Office hour (this week: Mol. Bio. Primer)
- Self-guided exercises to help you learn to use R for analysis.
- Using your own computer (other options possible)
- Exercise material will be made available ahead of time
- Towards end of course, more time devoted to working on group projects.

Readings

- No textbook, but we can give suggestions
- Lectures often come with suggested background papers (reviews or primary literature)
- Helpful to access journals online (e.g. via the UBC VPN)
 - http://it.ubc.ca/services/email-voice-internet/myvpn/setup-documents

Evaluation

- Homeworks
 - Two assignments worth 20 points each
 - +5 points each for peer evaluation
- Group project
 - Planning + project + poster session 40 points
- 10 Points for "other"
 - e.g. Preparedness, participation.

Homework assignment

- One for February, one for March.
- Involve detailed analysis of real data
- Deliverables include a short report and R code
- Two weeks from assignment to due date
- Lateness penalties

Group projects

- Starts today start thinking about it
- A few minutes for group project pitches on Jan 19 and Jan 21.
- Form groups by Fri Jan 23 (3-4 people)
- Friday Jan 30: initial project proposals
- Feedback to groups Feb 13 Proposals finalized by Feb ~15
- Work on projects over rest of term
- Final session of the course is the poster session

Group projects: where do they come from?

- Historically, almost all projects have been based on a data set provided by a student (i.e., collected in their lab).
- Occasionally, instead based on an idea from a student, using published data.
- If you need help thinking up an idea for a
 project let us know. But this has never been needed
 before (beyond refinement). If you are unsure of where you are
 going to get a project from, wait until you hear the project
 pitches.

Examples of past group projects

- · Genomic copy number alterations for prognosis of prostate cancer
- Learning about proteins from other proteins: Protein Database Prediction
- Conditional epistasis profiling in yeast
- Epigenetic biomarkers for cancer diagnosis
- Comparative metagenomics : metabolic potential
- · Epigenome and transcriptome in rice strains
- Analysis of HPV E2 protein on host gene expression
- Effects of Mutations in Histone Modifying Enzymes on Gene Expression Profiles
- Methodological considerations in analysis of Illumina Infinium methylation data
- Gene expression in invasive ragweeds
- Modeling time-course expression of SET domain-containing genes in mouse embryos
- · Gene expression in blood of humans with asthma challenged with allergen

2011 and 2012 project titles, paraphrased

High-dimensional biology

- I. What is it
- 2. What kinds of methods are used to analyze it
- 3. Some examples

Collecting data the lowdimensional way

- Pick one variable (e.g. "activity of a protein") and study it under various conditions.
- Repeat this for another variable
- Usually "hypothesis-driven"
- Powerful, but knowledge accumulates slowly and synthesis is difficult

Biology is complicated

- Thousands of "parts"
- Limitations of the "one thing at a time approach" how do the parts work together?
- Technology enabling increasingly detailed analyses – measure many things in parallel
- Drawback: Fishing expeditions?

Defining "high dimensional"

- Large number of features measured in each sample/subject/individual ("high content")
 - Genes, proteins, DNA sites, brain regions, etc.
- Not usually talking about huge numbers of samples (e.g. individuals studied) –
 - often 10s, but can be 1000s (some genetics studies)
- Studies can sometimes be "non-hypothesis driven"

Example of a question answered with a high-dimensional approach

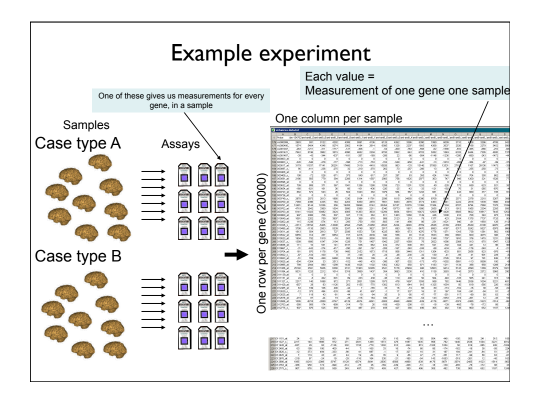
- Tumor type A is deadly and type B is more easily treatable (but still bad)
- Telling A from B is difficult
 - Cells look the same, etc. we only find out by seeing what happens to the patients.
- We know that cancer is a "gene" disease

Questions:

- Where is the difference?
- Can we find new targets for drugs or for diagnosis?
 - (Drug targets are usually proteins, encoded by genes)

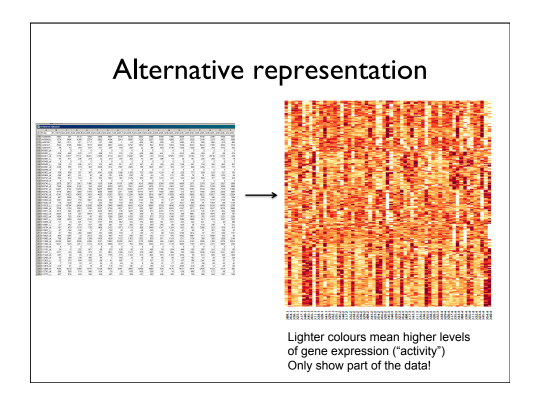
Looking for insight from genomics

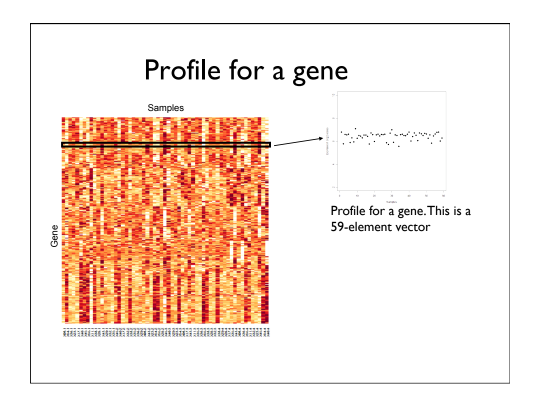
- Since cancer is a disease of genes, let's look at the genes not just one, but all of them
- We are hypothesizing that there is some difference in genes between the two types, if only we could find it
- But we're not starting with a specific hypothesis. We're going to test thousands of hypotheses
- In this example, we're going to look at "gene expression levels" a measure of "how active" is each gene.

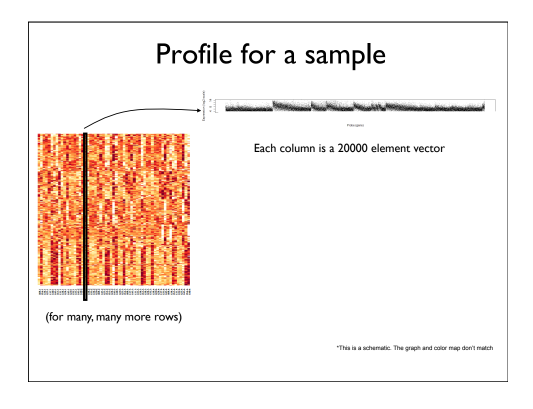


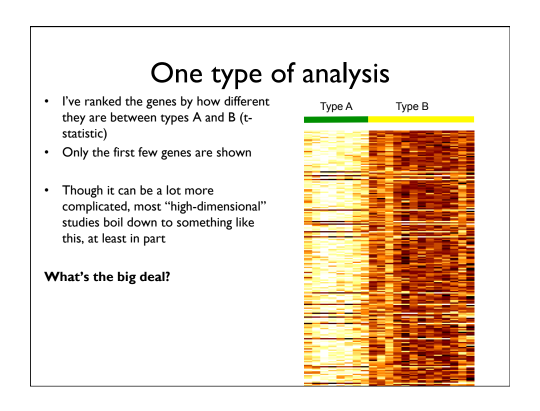
A partial list of things to assay

- DNA/Chromatin
 - Genotypes, copy numbers ("mutations" and variants)
 - DNA methylation
 - Chromatin state (histone marks, transcription factors ...)
- RNA
 - Quantification of transcripts (protein coding, non-coding)
 - Transcript variants (splicing, editing)
- Proteins
 - Detection, Quantification
 - Binding and complexes
- · Metabolites and other small molecules
- Phenotypic screens
 - RNAi (etc.)
 - Genetic interactions
- Cellular composition of a sample (cell types)







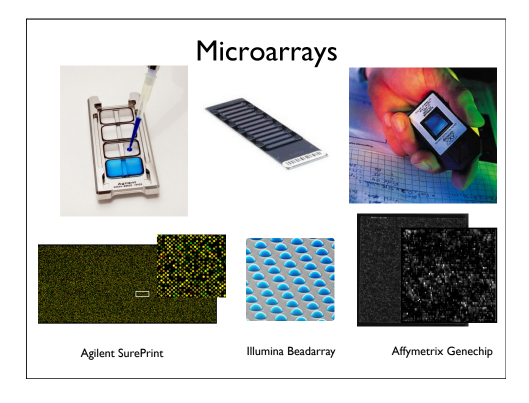


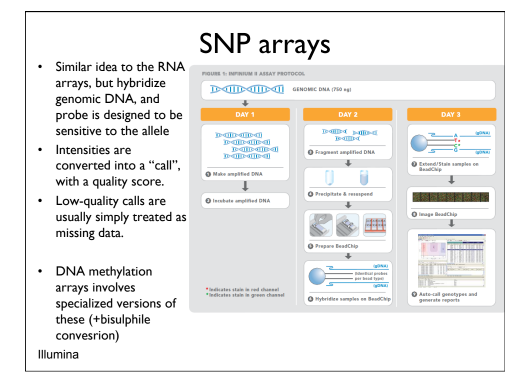
Pitfalls and challenges

- Signals can be small and buried in lots of nonsignals; False positives are a danger.
- Need to detect outliers, batch effects and other confounds
- Can we make better use of the fact that we're testing 20,000 genes than just doing a t-test on each one?
- Data sets (and questions) are often much more complex
- Getting just a list of "hits" isn't enough can we understand something more about the "system"

High-dimensional technologies

- DNA & RNA sequencing
 - Transcriptomes, exomes, full genomes
- Complex gene library construction
 - Expression vectors, protein tags, knockdowns
- Microarrays and other robotic/parallel tech.
 - Screens, high-content assays ...
- Mass spectroscopy
- Flow cytometry
- Imaging





Sequencing-based assays

- Instead of using hybridization to a designed probe, determine the DNA sequence of the sample
- Several competing platforms
- Genotyping: Compare to a reference
- RNA: quantify how many times you see a sequence





Illumina HiSeq

Analysis modes

- What is the general toolkit available for the analysis of data?
- How are these specialized for highdimensional data?

Exploratory analysis

- The first thing you do with your data
- Graphs and other visualizations, often combined with data reduction
- Use to spot problems, formulate hypotheses
- Often rely on power of human brain
- Data reduction essential to make exploration tractable for large data sets, even then it can be a challenge
- Follow up with more formal analysis

Model fitting and hypothesis testing

- Formally test a specific question about the data
- Is what I see "statistically significant"?
- False positives are a major risk in large data sets
- Can exploit repeating structure of the data to improve ability to find true positives

Unsupervised learning

- "Learn" undiscovered groupings in the data
- Clustering -- how do my samples or features group together?
- Useful as an exploratory technique as well as "data mining" when backed with quantitative analyses
- Example: Finding previously unknown groups of subjects based on a gene profile

Supervised learning

- Can I predict an unmeasured feature of a sample from a measured one?
- Less common than unsupervised learning, most commonly used in clinically-oriented settings – development of biomarkers
- Example: predicting tumour drug response based on gene profiles

Other methods

- Many analyses just give a list of genes
- "Downstream" analysis needed to make sense of it "biological interpretation"
 - Overlay/combine/compare with other data
 - Transform one data set into another type of data at a different granularity
 - Genes → pathways
- Usually these end up returning to exploratory etc. modes

More examples

- Illustrate some real-life cases of highdimensional data
- We hope to teach you enough in the course to do at least primitive versions of these analyses
- ... or at least be able to read the papers
- ... even if it's a type of experiment we don't teach in detail.

