

Bioinformatics: from sequence to structure Module 2

Statistical machine learning

Devika Subramanian

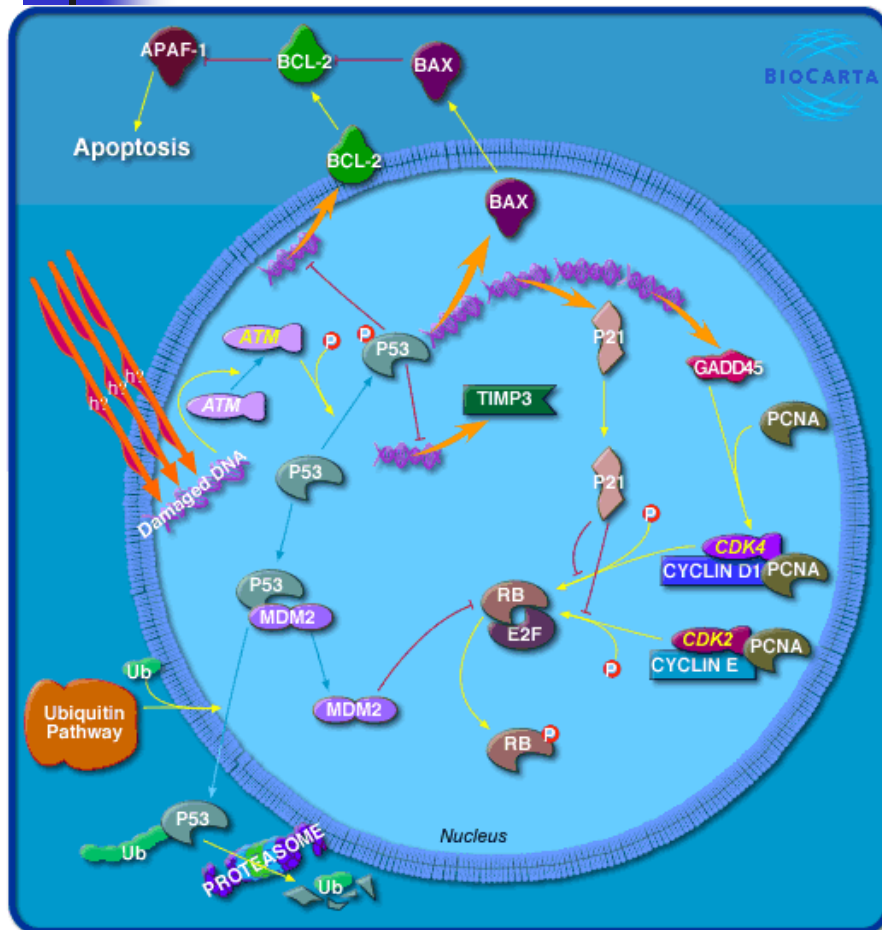
Comp 470



Module design inspiration

- "...Deciphering how a mere 10^7 nucleotides result in a yeast cell, let alone how 3×10^9 nucleotides result in a human - cannot begin until the **genes have been annotated**. This step includes figuring out **the proteins these genes encode and what they do for a living**. But understanding **how all of these proteins collaborate to carry out cellular processes** is the real enterprise at hand."
 - -- ----- Stanley Fields (Science:Feb 16 2001: 1221-1224)

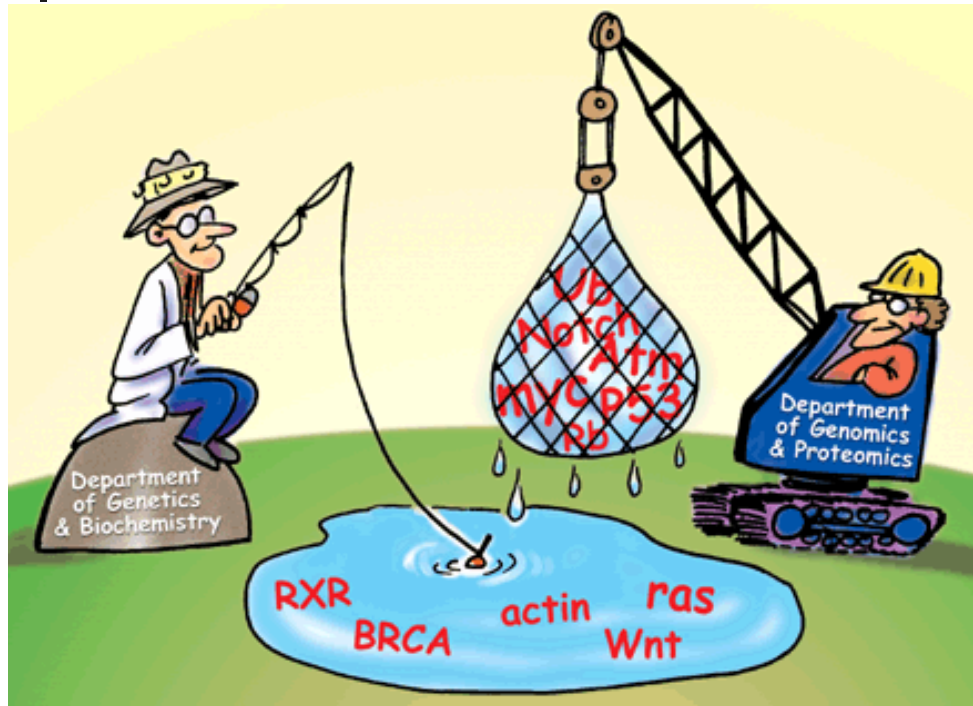
Signaling & metabolic networks



- ◆ Consist of interacting proteins, genes and, small molecules.
- ◆ Underlie the major functions of living cells.

The quest:
The wiring diagrams
of life, particularly how
they are altered in
diseased cells.

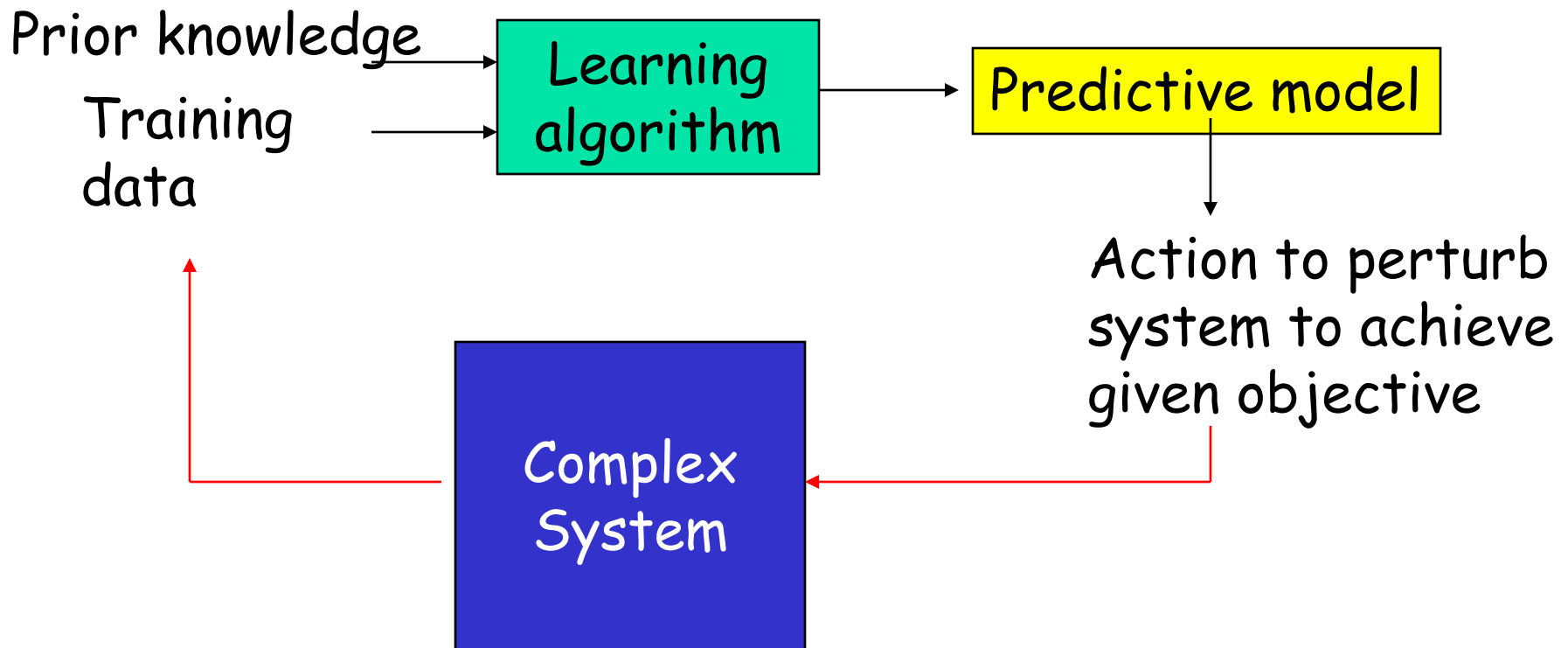
Building models from data



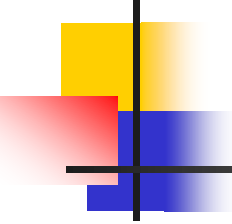
- 3 billion base pairs in human genome.
- 1.5 million known proteins.
- 10^6 to 10^9 (projected) protein-protein interactions.

High throughput assays: mRNA expression levels of 15,000 genes in 1 shot, flow cytometry, SELDI-TOF proteomics assays, allow us access to cellular processes

What is machine learning?



Observe complex system and build models to predict its response.



Fundamental questions in machine learning

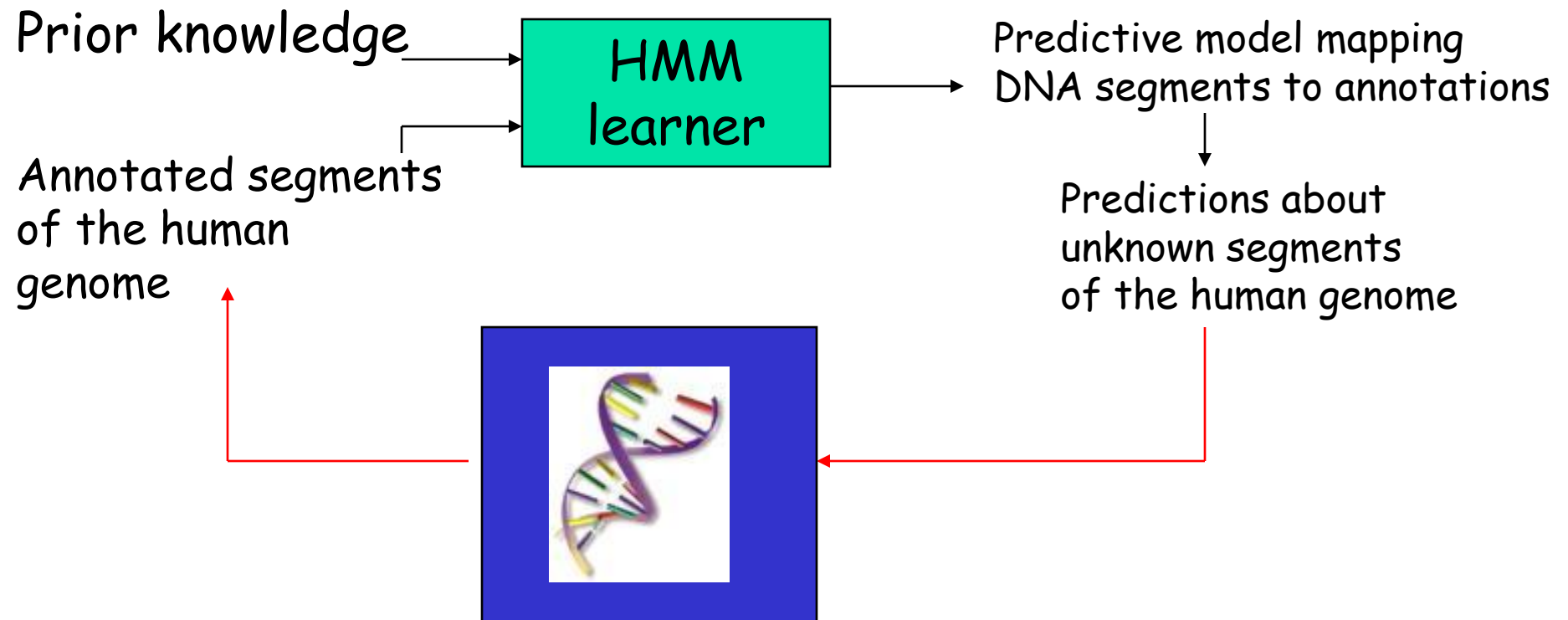
- What aspects of the system to observe? (Feature selection)
- What class of models to build from observed data and prior knowledge? (Model selection)
- How to evaluate efficacy of the learned model? (Model validation)



Three illustrative problems

- Given a DNA sequence, find and annotate genes in it.
- Given gene expression data, determine biologically significant genes that are differentially expressed.
- Given flow cytometry data, learn signaling networks in normal and diseased cells.

Computational Genefinding

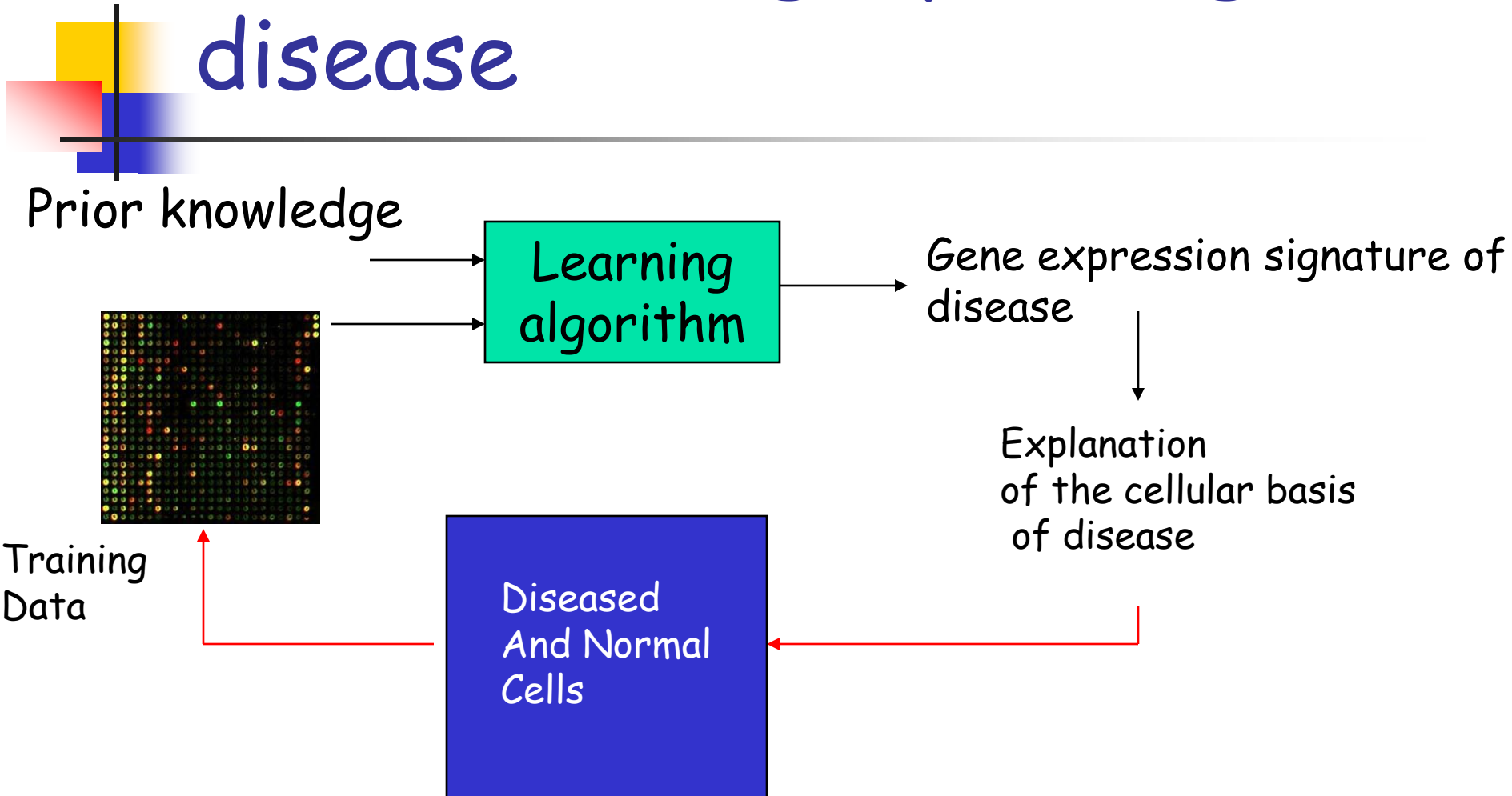




Three illustrative problems

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Molecular fingerprinting of disease

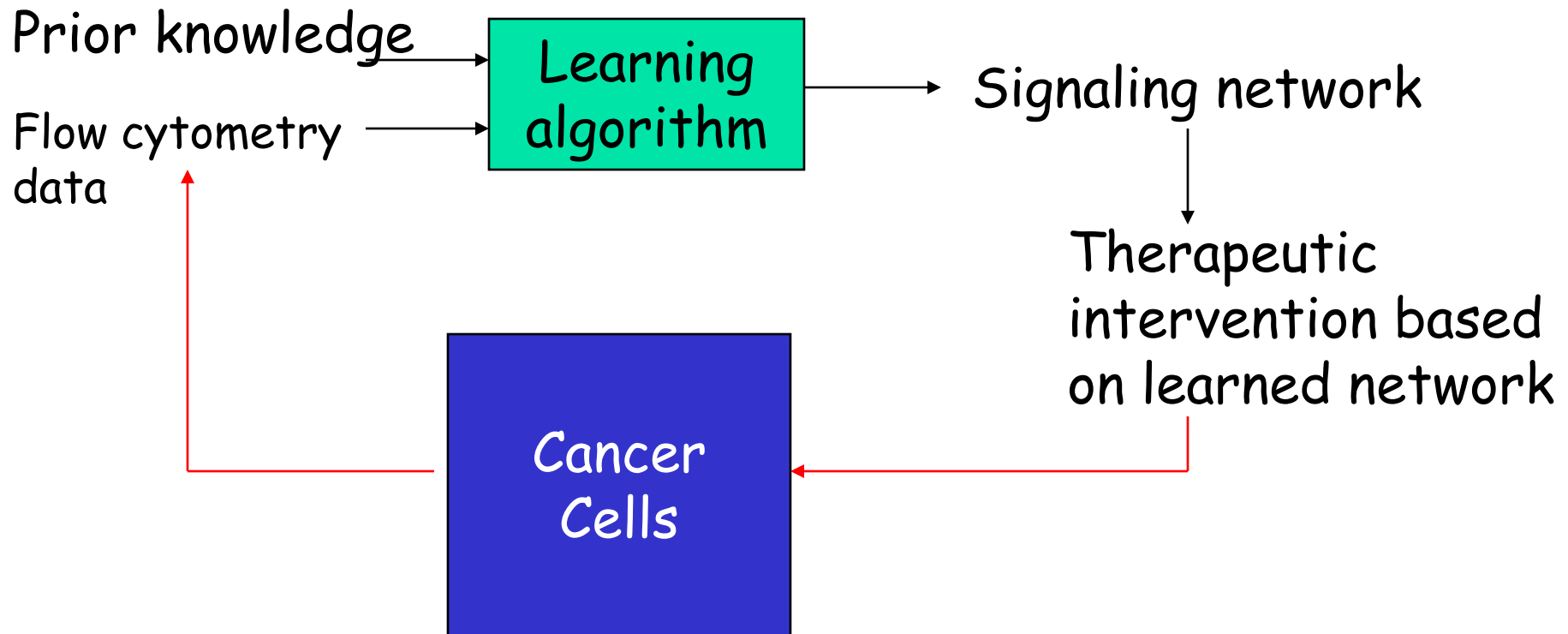




Three illustrative problems

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Learning cell signaling networks from data





Three statistical learning algorithms

- Hidden Markov Models and variants.
- Naïve Bayes classifiers and support vector machines.
- Bayesian network learning: parameter and structure learning.

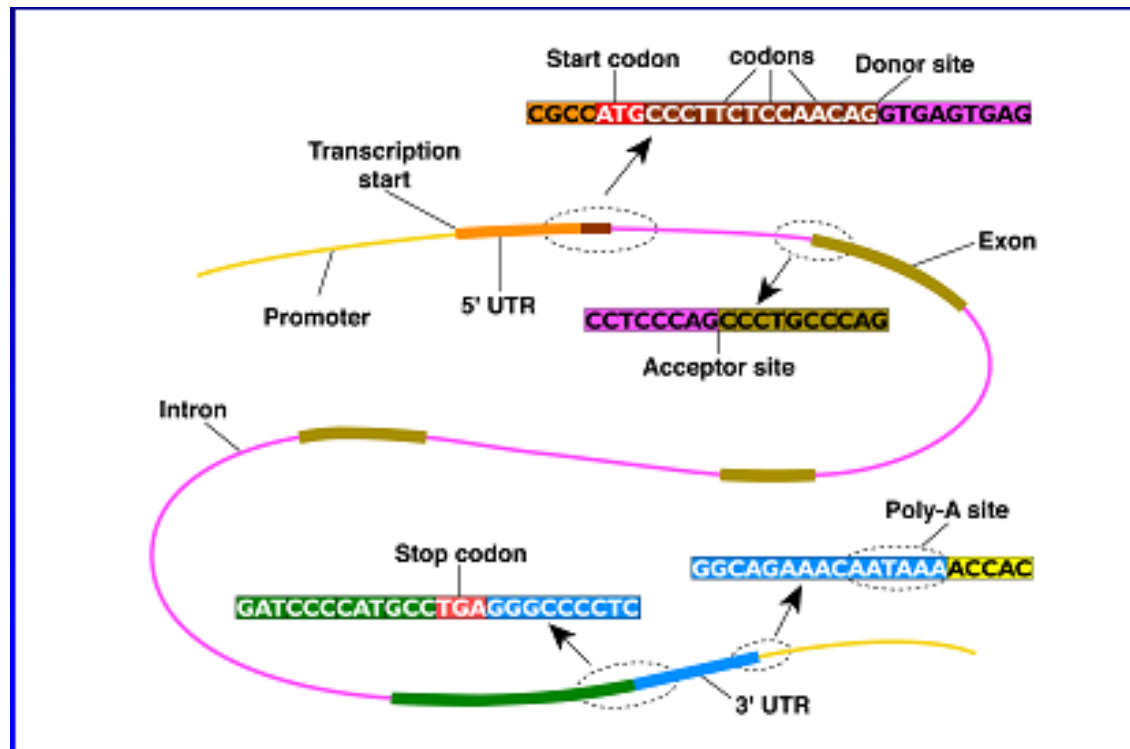


Module objectives

- Learn to model heterogeneous biological data and choose appropriate statistical machine learning algorithms.
- Understand the basics of supervised and sequential machine learning algorithms with particular focus on hidden Markov models, naïve Bayes classifiers, kernel-based methods and Bayesian networks.
- Apply these techniques in the context of real data (human chromosome 22, prostate cancer gene expression data, flow cytometry data from T-cell signaling).

Computational gene finding

- Gene finding in eukaryotic DNA



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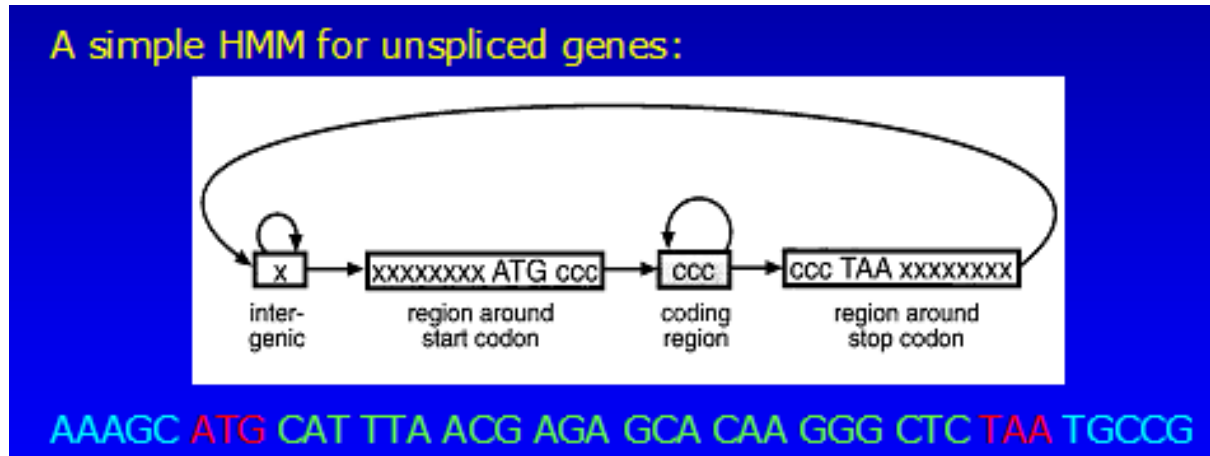


Mathematical model

- Hidden Markov models
 - Structure of HMMs
 - Viterbi algorithm for annotation
 - Baum-Welch (EM) algorithm for learning models
 - Extensions: pair HMMs

Ab initio methods

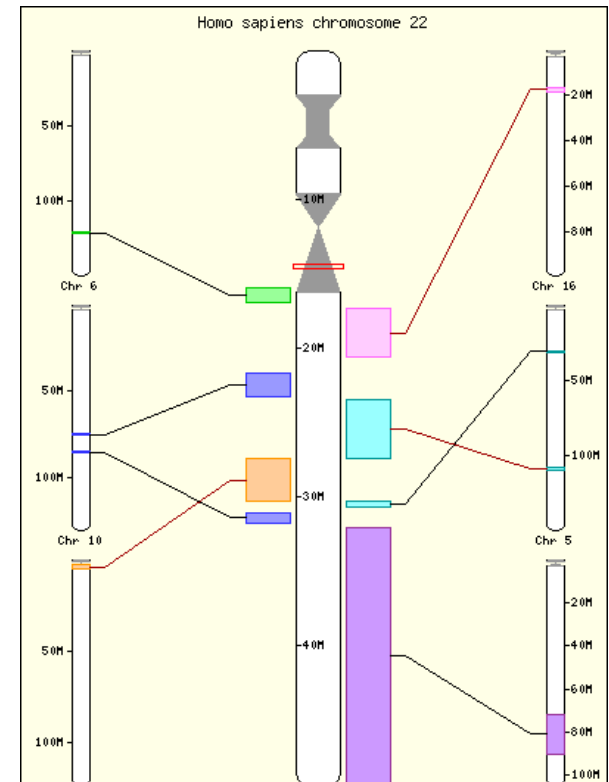
- Genscan (Burge et. al., JMB 1997)



- Intrinsic limits on performance of *ab initio* methods; evaluation studies (Rogic, *Gen. Res.* 2001)

Comparative methods

- SLAM (Pachter et. al. *Gen. Res.* 2003); simultaneous gene prediction and sequence alignment of two syntenic genomic regions.
- Paired HMMs



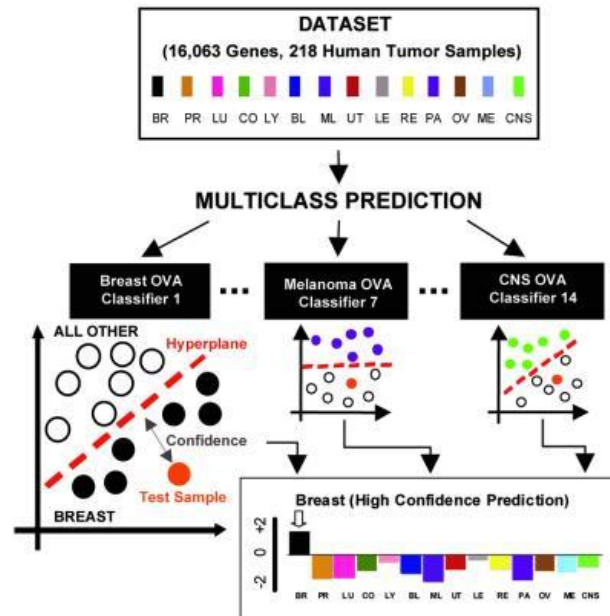


Exercise

- CpG island detection on human chromosome 22 using learned HMMs.
- Analyze similarities and differences in prediction between Viterbi and posterior decoding.

Molecular fingerprinting of cancers

- Work of Golub et. al (Science 1999 (AML/ALL), Bioinformatics 2001, Nature 2003), Lee & Lee (Bioinformatics 2003)





Mathematical model

- Naive Bayes classifiers
 - Ensemble methods: boosting and bagging
- Support vector machines (SVM)
 - Maximum margin separating hyperplane
 - Linear SVMs and soft margin hyperplanes
 - Non-linear SVMs and the kernel trick



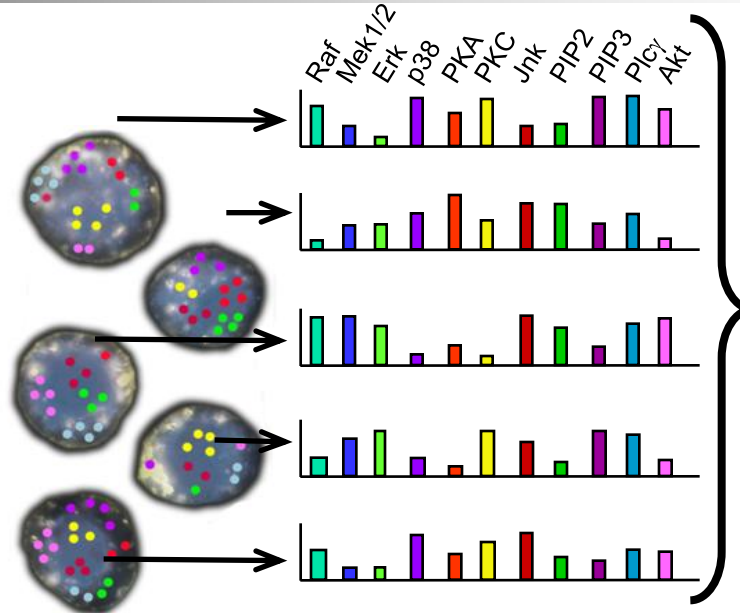
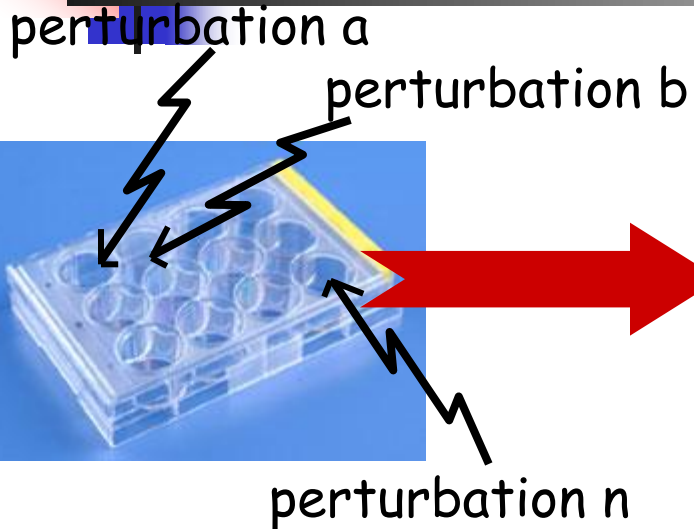
Exercise

- From Singh prostate cancer data, determine which genes are differentially expressed using Naïve Bayes and SVM classifiers.
- Experiment with various feature selection techniques, compare predictions against the latest theories of compromised cellular processes in prostate cancer (Science 2004).

Learning networks from data

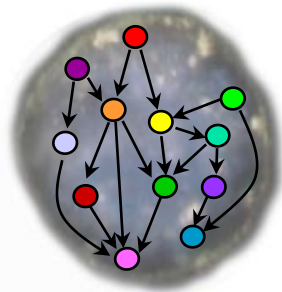
Conditions (96 well format)

Multiparameter Flow Cytometry



Datasets of cells

- condition 'a'
- condition 'b'
- condition... 'n'



Influence diagram of measured variables

Bayesian Network Analysis

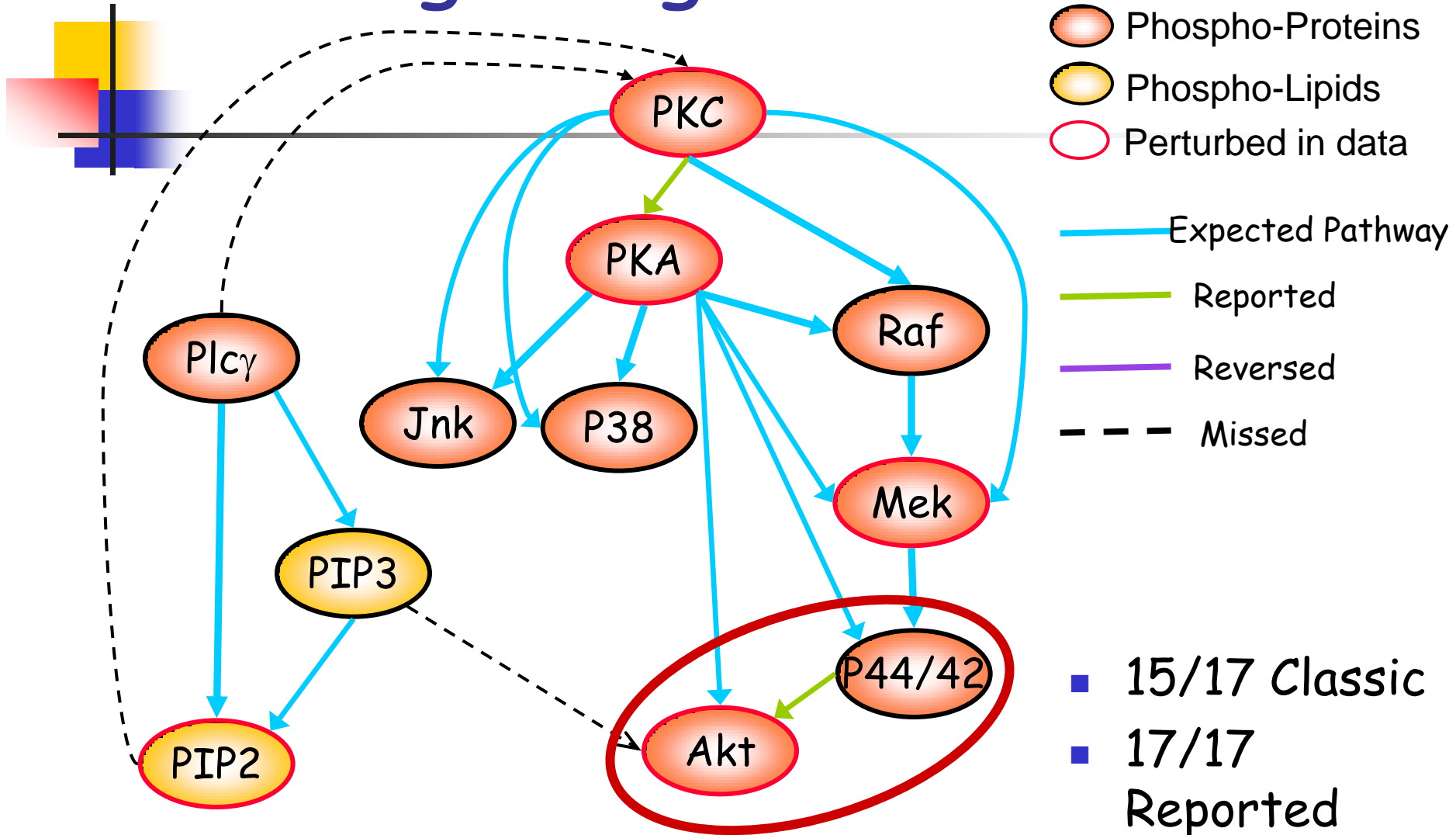
, Devika Subramanian, 2009



Mathematical model

- Probabilistic models : bayesian network representations of signaling networks.
- The sparse candidate algorithm for learning Bayesian networks from high-throughput data .

T-cell signaling network



Science 2005, Sachs et. al

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- 15/17 Classic
- 17/17 Reported
- 3 Missed



Summary

- How to use the underlying biology to constrain model selection and feature selection.
- How to choose and adapt machine learning algorithms for biological problems.
- How to design learning protocols to deal with incomplete, noisy data.
- How to interpret the results of machine learning algorithms.