Exploration Session Week 8: Computational Biology

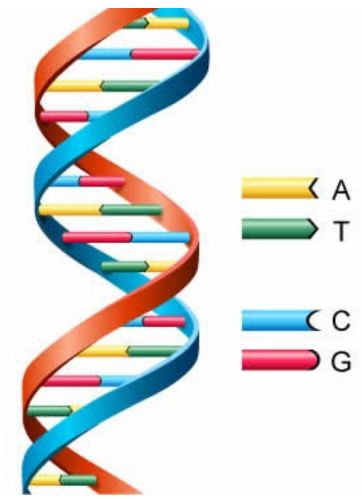
Melissa Winstanley: mwinst@cs.washington.edu

(based on slides by Martin Tompa, Luca Cardelli)

Exploring DNA Sequences

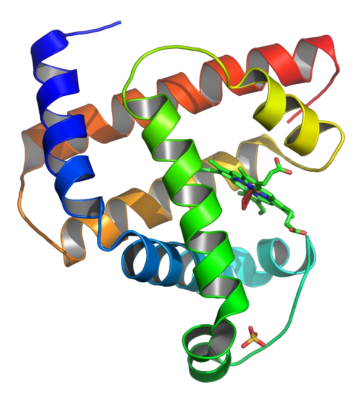
Overview of DNA

- Instructions for cellular functionBuilding proteins
- Composed of *nucleotides* Adenine, thymine, cytosine, guanine
 A pairs with T, C pairs with G
- Double-stranded: forms a double helix
 - Strands have an orientation
 - Pairing of antiparallel strands
- Huge amount of DNA
 - □ 3 billion base pairs, 2m long in a cell
 - 133 AU long in human
 - **2**0 million light years long in human population



Overview of Proteins

- Workhorses of cells
- Composed of sequence of *amino acids* 20 to 5000 amino acids in a protein
- 20 possible amino acids
- Proteins fold into complex 3D shapes
 - Fold-It
- Information to make proteins encoded in DNA
 - Codon: 3 base pairs
 - Ex. CTA \rightarrow leucine
 - Gene: sequence of DNA for 1 protein



Overall Goals

Overall

- Identify key molecules in organisms
- Identify interactions among molecules
- Computational focus: sequence analysis
 - Identify genes
 - Determine gene function (what protein is produced?)
 - Identify proteins involved in gene expression
 - Identify key functional regions
- Why do we care?
 - Determining function of a new sequence
 - Genetic diseases
 - Evolution

String Alignment

How to judge how well two strings are aligned?

acbcdb a c - b c d b cadbd - c a d b - d -

- Each dash represents an inserted space
- Assign +2 to every exact match, -1 to every mismatch

3 * 2 + 5 * (-1) = 1

Higher score indicates a greater match between the strings

BLAST Algorithm

- "Basic Local Alignment Search Tool"
- For comparing biological sequence information
 - Amino acid sequences (proteins) or nucleotide sequences (DNA)
- Inputs
 - A query sequence Q
 - A database D of sequences
- Output
 - Sequences from D that match Q above a certain threshold
- Usefulness
 - Unknown gene in a mouse, so query the human gene database to see if a similar gene exists in humans

Make k-letter subsequences from Q Ex. k = 3: "acbcdb" "acb' "cbc" "bcd" "cdb"

Usually k = 28 for DNA, k = 3 for proteins

□ For each subsequence w, find matching subsequences

- Only consider a matching subsequence if its alignment score is greater than some threshold
- Alignment(seq) >= T

seq = "ACT" \rightarrow Alignment = 2 * 1 + 2 * (-1) = 0 Not considered

- Scan the database for exact matches with the high scoring subsequences
- Take each exact match and extend in either direction (no gaps)
 - Until the score decreases below a "dropoff"
 - Forms a "high-scoring segment pair" (HSP)
- Only save match extensions above a certain score threshold S

HSP: score = 2 + 2 + 2 - 1 + 2 = 7

□ For each HSP, do a gapped extension (spaces possible)

Output each extension that has probability of randomly occurring below a pre-set threshold x

More Complicated Analysis

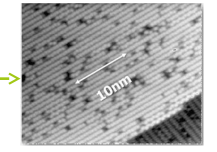
- Multiple sequence alignment
- Different ways to score subsequences
- Considering context around a sequence
- Predicting 3D structures of proteins

Programming Molecules

Getting Smaller

First transistor

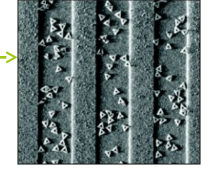
25nm NAND flash

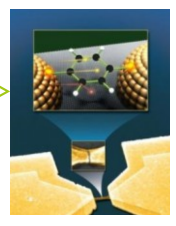




Single molecule transistor

Molecules on a chip





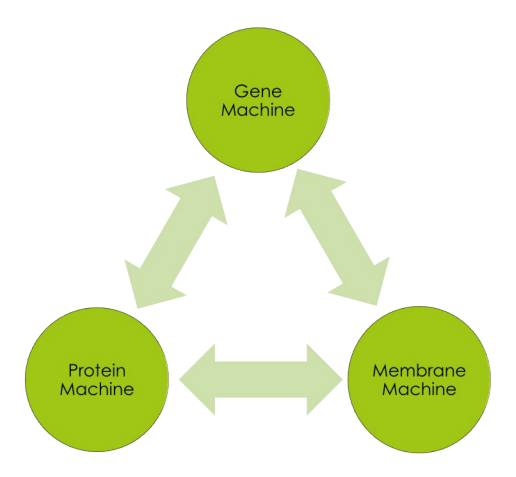
~10 Moore's Law cycles left

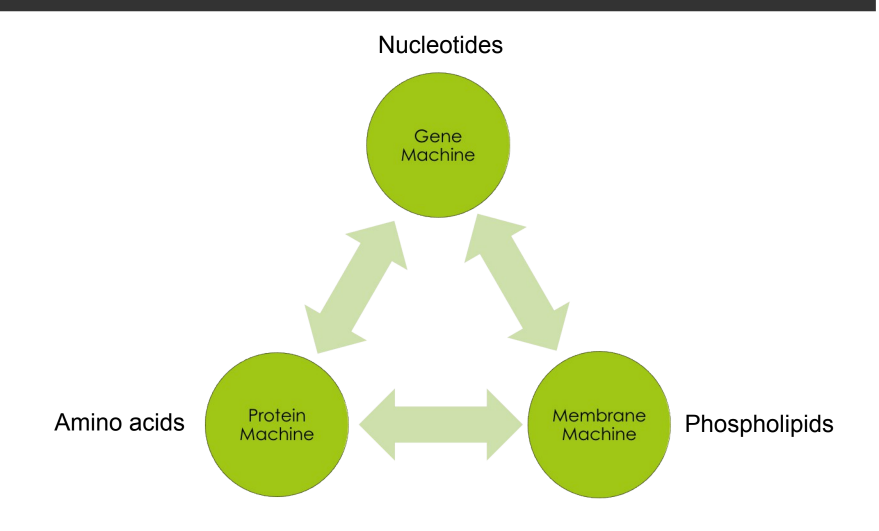
http://upload.wikimedia.org/wikipedia/commons/thumb/b/bf/Replica-of-first-transistor.jpg/200px-Replica-of-first-transistor.jpg http://www.blogcdn.com/www.engadget.com/media/2010/01/01-30-10intelflash.jpg http://www.wired.com/images_blogs/gadgetlab/2009/12/molecular-transistor-264x300.jpg http://www.internetnews.com/img/2009/08/ibm_dna_chips.jpg

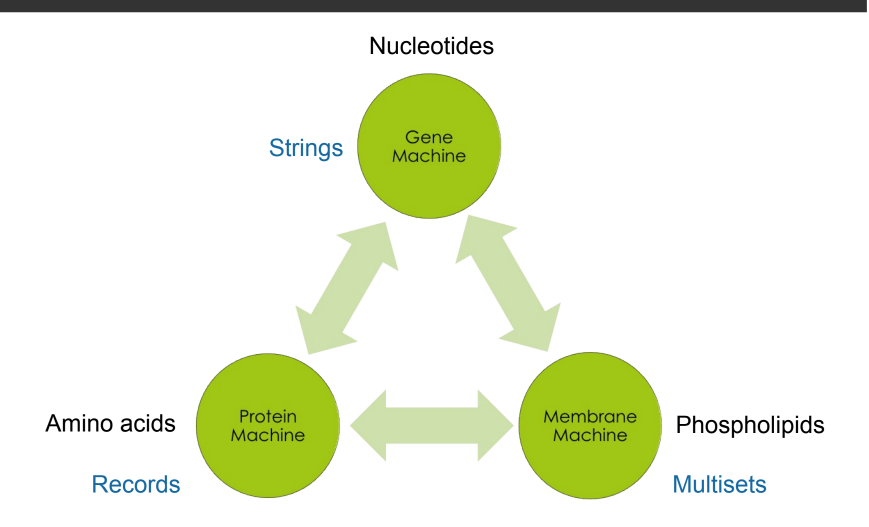
Building Smaller

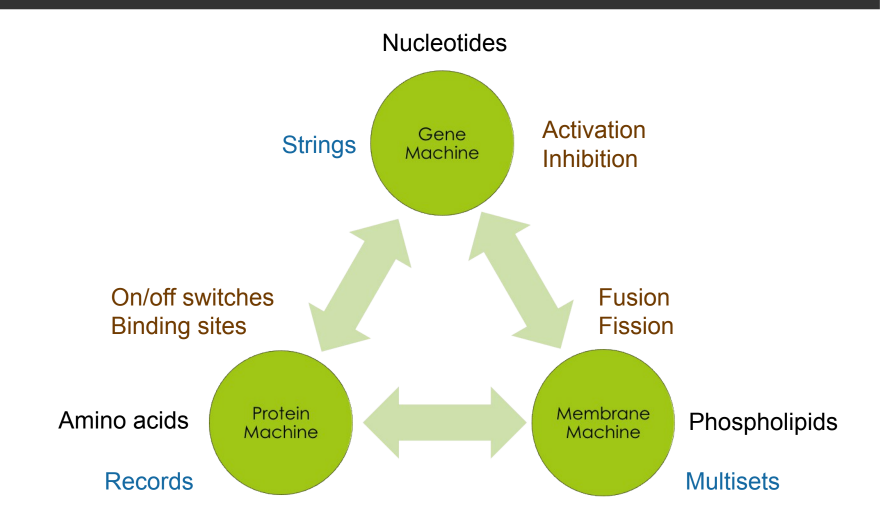
- How to build things smaller than your tools?
- You can't
 - Solution: self-assembly
- Molecular IKEA
 - Dear IKEA, please send me a chest of drawers that assembles itself.
- At a molecular scale, many such materials exist
 - Proteins, DNA/RNA, membranes
 - http://youtu.be/0N09BIEzDII









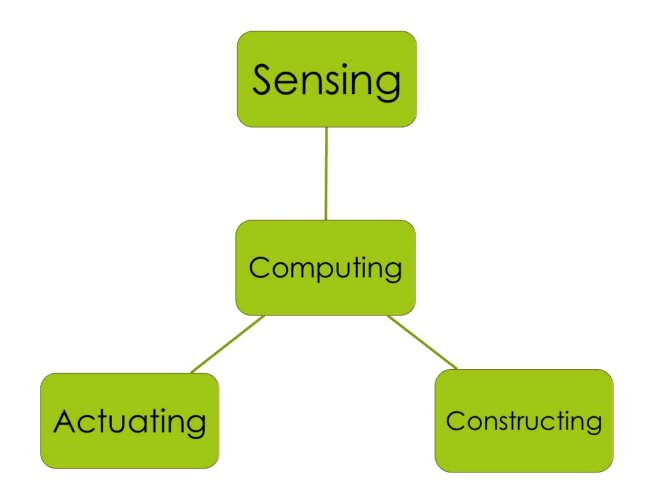


How do we form a "language"?

- Chemical reactions
 - $\Box A + C \rightarrow_{\Gamma} B + D$
 - Instructions in a "program"
- Problem: combinatorial explosion
 - SO MANY chemical reactions in a cell
- Model reactions as automata machines that perform a task

- Problem: chemistry is not an executable language
 - Dear Chemist, please execute this arbitrary reaction.

Controlling Systems on a Nanoscale



DNA Tweezers



One Approach to Autonomous Computing

- Goal: precisely control organization and dynamics of matter and information at the molecular level
 - Uses DNA, but use is accidental
 - No genes involved



"Gates" and "transducers"

Molecular programming workflow

- First figure out what gates you want to use and signals you want to send
- □ Signals + gates \rightarrow structures of DNA
- □ Structures \rightarrow sequences of DNA (NUPACK)
- □ Sequences \rightarrow DNA synthesis (IDT)
- DNA synthesis \rightarrow mail
- □ Receipt of DNA \rightarrow_{water} execution
- Florescence is your "print" statement