### Genome 540

# Introduction to Computational Molecular Biology:

Genome and protein sequence analysis

## Today's Lecture

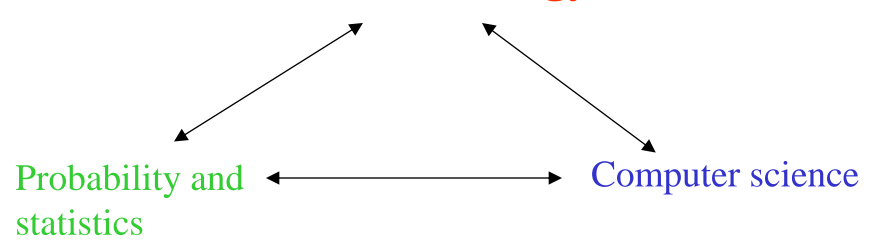
Course overview

Administrative details

• Finding exact matches in sequences using suffix arrays

## Computational Molecular Biology

## Molecular biology



## Course Lecture Content

DNA and protein sequences

- Algorithms
  - Dynamic programming

- Probability models
  - HMMs

#### We do *not* cover:

- 'Non-linear' (non-sequence based) computational biology
  - protein structure, expression arrays, metabolic pathways, models for interacting molecules ...
- 'Machine learning' applications
- Existing software tools

## Course Prerequisites

- You *must* 
  - be able to write programs for data analysis
  - have access to a computer where you can write & run your programs

HW assignment # 1 will be a good test!

- Some previous familiarity with
  - probability and statistics
  - molecular biology
  - is highly desirable
    - (if you lack it, you will have to work harder!)

## Course Requirements

- Homework
- No tests or exams
- Attendance at discussion section strongly encouraged but not required
- Ask questions!
  - in lecture
  - at discussion section
  - by email
  - (via message board)

### Homework

- Due weekly, Sunday at midnight
  - Posted on web site approx 1.5 weeks in advance
  - Each is 10 pts, late penalty of 1 pt/day (max penalty 3 pts)
  - Can redo
- write computer program to analyze genomic data set
  - "From scratch", i.e. not using prewritten routines from elsewhere
  - Run on your own computer
  - Programming language is up to you but a compiled language
    (e.g. C, C++) is recommended for efficiency reasons
    - Python + Cython also works
    - Interpreted language may work, but risky!

- Also: readings (in textbooks, or journal articles)
- turn in results of analysis, and your program, with (in some cases) a written interpretation of the results;
  - all to be submitted by email in computerreadable format

### Course Info

- Instructors (contact info is on web page):
  - Phil Green
  - TA: Mitchell Vollger

- Office hours by appointment (send Mitchell or me an email)
- if you did not receive the email I sent yesterday, send me (<a href="mailto:phg@uw.edu">phg@uw.edu</a>) your email address today (whether or not you are registered!)

- Lectures: TuTh 10:30-11:50, Foege S-110
- Weekly discussion section:
  - discuss homework, answer questions
  - review background material
  - related topics (next-gen sequencing?)

#### Tentative time/place: Th 12-1, Foege S-040

- If you have a conflict, email me your schedule of unavailable times -- we may be able to find another
- Web site: <a href="http://www.phrap.org/compbio/mbt599">http://www.phrap.org/compbio/mbt599</a>
  - will post HW assignments, copies of lecture & discussion section slides here
  - has link to last year's site for approx syllabus & slides

## Texts (will follow only loosely):

- Biological Sequence Analysis: Probabilistic Models of Proteins and Nucleic Acids by Durbin, Eddy, Krogh & Mitchison. Paperback, ~\$60.
- Statistical Methods in Bioinformatics: An Introduction (Statistics for Biology and Health) by Ewens & Grant. Hardbound, ~\$105. **N.B.** This is the **2D edition!**
- available from UW Bookstore (South Campus Center branch) or from Amazon or Barnes & Noble

# Finding perfectly matching subsequences of a sequence

- Idea (*much* more efficient than 'brute force' approach):
  - suffix array (Manber & Myers, 1990)
  - make list of pointers to all positions in sequence
  - lexicographically sort list of strings that are pointed to
  - process the list: adjacent entries are "maximally agreeing"

## Suffix array step 1: List of Pointers to Suffixes

#### ACCTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC

ACCTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_1$ CCTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_2$ CTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_3$ TGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_4$ GCACTAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_5$ CACTAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_6$ ACTAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_7$ CTAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_8$ TAAACCGTACACTGGGTTCAAGAGATTTCCC  $p_9$ AAACCGTACACTGGGTTCAAGAGATTTCCC  $p_{10}$ AACCGTACACTGGGTTCAAGAGATTTCCC  $p_{11}$ ACCGTACACTGGGTTCAAGAGATTTCCC  $p_{12}$ 

# Suffix array step 2: View as Strings to be Compared

#### ACCTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC

$p_1$	ACCTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_2^-$	CCTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_3^-$	CTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_4$	TGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_5$	GCACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_6$	CACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_7$	ACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_8$	CTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_9$	TAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_{10}$	AAACCGTACACTGGGTTCAAGAGATTTCCC
$p_{11}$	AACCGTACACTGGGTTCAAGAGATTTCCC
$p_{12}$	ACCGTACACTGGGTTCAAGAGATTTCCC

## Suffix array step 3: Sort the Pointers Lexicographically

#### ACCTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC

$p_{10}$	AAACCGTACACTGGGTTCAAGAGATTTCCC
$p_{11}^{-1}$	AACCGTACACTGGGTTCAAGAGATTTCCC
$p_{28}^{}$	AAGAGATTTCCC
p <sub>17</sub>	ACACTGGGTTCAAGAGATTTCCC
$p_{12}^{-1}$	ACCGTACACTGGGTTCAAGAGATTTCCC
$p_1$	ACCTGCACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_7$	ACTAAACCGTACACTGGGTTCAAGAGATTTCCC
$p_{19}$	ACTGGGTTCAAGAGATTTCCC
<b>p</b> <sub>29</sub>	AGAGATTTCCC
p <sub>31</sub>	AGATTTCCC
p <sub>33</sub>	ATTTCCC
p <sub>27</sub>	CAAGAGATTTCCC
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