

Hands-on Session 2: Obtaining Data from On-line Sources

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Session Organization

- **Goal:** To be comfortable building trees from real data
- **Lecture:**
 - Standard Software Packages
 - Details on Web-based Software
 - Motivating Problem
- **Lab:**
 - Organized so you can use the DIMACS lab, or your own laptop
 - Welcome to work singly or in groups

Lecture Outline

- Motivating Problem

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- Building Trees Overview

Lecture Outline

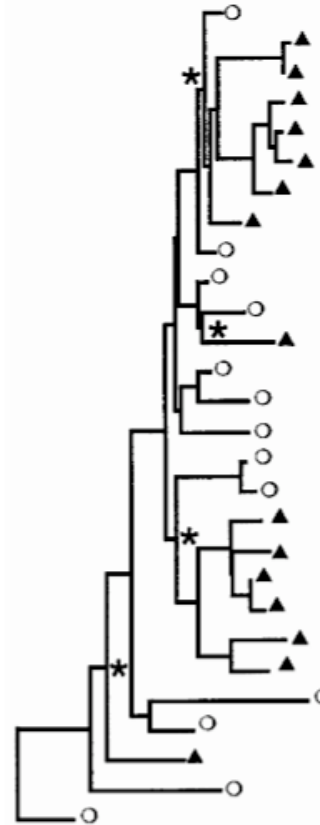
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- Using Sequence Databases

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- Using Sequence Databases
- Aligning Sequences

Motivating Problem: Building Trees with Serial Data?

Rodrigo *et al.*,
“Coalescent estimates of
HIV-1 generation time in vivo.”
PNAS '99



Motivating Problem: Using Serial Data

- Rodrigo *et al.* includes 55 HIV-env partial sequences, all from the same patient

Table 1. Summary statistics for each sequence sample set

Sample	Days from first sample	No. of sequences	Average pairwise diversity, %	θ	N
1	0	13	3.6	0.088	1100
2	214	15	3.9	0.106	1325
3	671	15	5.0	0.074	925
4	699	9	4.2	0.144	1800
5	1005	8	4.1	0.092	1150

- Starting question: what is the genealogy samples (from the same patient) taken at different times?

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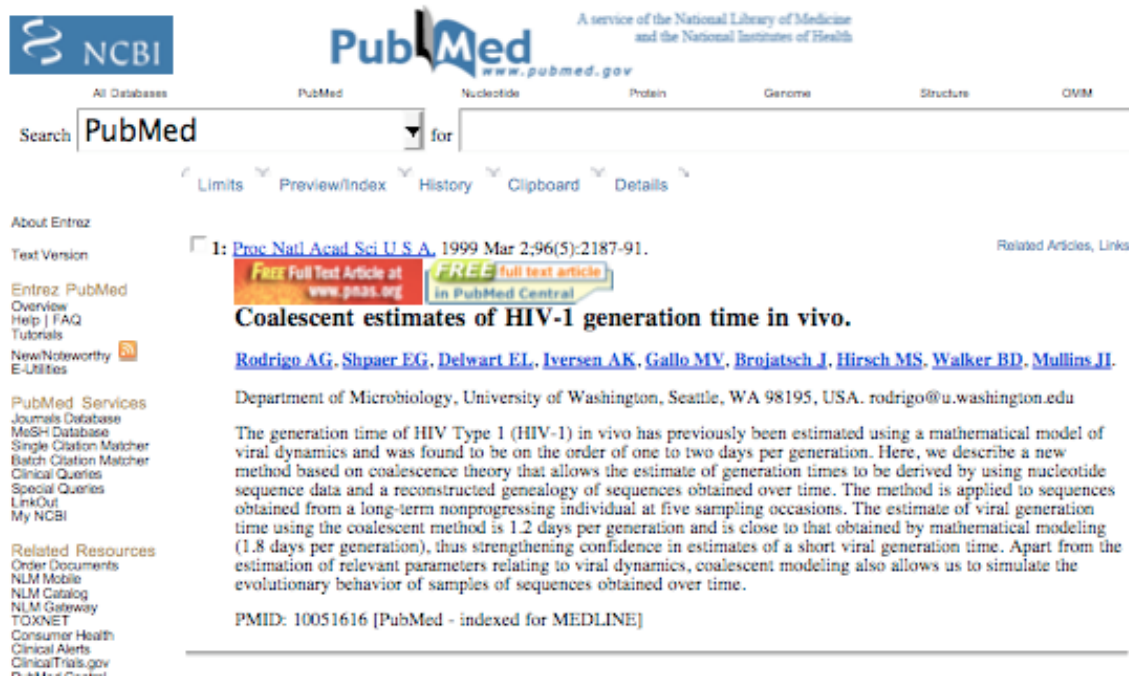
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We'll focus on the first two today.

Using PubMed

An on-line index of scientific papers:



Can search by all standard methods...

Sequence Databases

- GenBank: repository of sequences from NCBI (NIH).
- As of August 2005, GenBank had 100 gigabases of sequences.
- Almost all sequences from published articles are there, and can be located by their unique **accession number** or PubMed ID.

LANL HIV Databases

- Los Alamos National Laboratory maintains databases of sequences, resistance, immunology, and vaccine trials.
- Can be searched in numerous ways including accession number or PubMed ID.

Aligning Sequences

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- One of the most common alignment programs is ClustalW:
 - Available via multiple servers including EBI & the Pasteur Institute
 - Does a global multiple sequence alignment

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- Using your favorite phylogenetic reconstruction method, build a tree from the sequences.
- Analyze resulting trees

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For example, after returning the distance matrix, you have the option of applying a method to the matrix.

Helpful Websites

- Dataset for this tutorial:

<http://comet.lehman.cuny.edu/stjohn/dimacsTutorial>

- PubMed & Genbank:

<http://www.ncbi.nlm.nih.gov/entrez>

- HIV Sequence Database:

<http://hiv-web.lanl.gov/content/index>

- The Pasteur Institute:

<http://bioweb.pasteur.fr/intro-uk.html>