



Introduction to molecular epidemiology and infectious disease phylodynamics

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SISMID, July 19-21, 2017

This course (SISMID module 13)

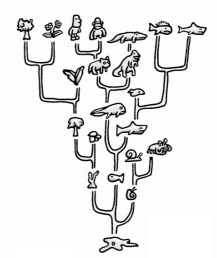
- Wednesday, July 19
 - → Introduction
 - → Alignment, substitution models and phylogenetic inference
- Thursday, July 20
 - → Phylogenetic inference practical
 - → Bayesian phylogenetics
 - → Molecular clocks and model testing
 - **⇒** BEAST practical
 - http://rega.kuleuven.be/cev/ecv/

- Friday, July 21
 - → Viral epidemiology and the coalescent
 - → BEAST practical
 - → Phylogeography
 - **⇒** BEAST practical
 - Bonus
 - → Phylo-Alignment
 - → Recombination
 - → Robust Counting
 - → Antigenic cartography

(We are here to cater for your needs!)

Molecular evolution and phylogenetics

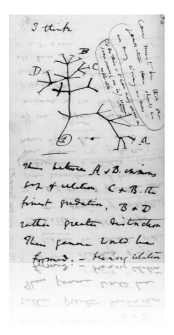
- biological sequences (DNA, RNA, protein) contain information about the processes and events that formed them
- this information is often scrambled, fragmentary, hidden, or lost completely
- our aim is to use mathematical models to recover and decipher this information
- The central concept is a phylogeny: a diagram depicting the ancestral relationships among characters or genetic sequences



HIV-1 (UK) ATC---TGCTAAAGCATATGACACAGAGGTACATAATGTTT
HIV-1 (USA) ATCGGATGCTAGAGCTTATGATACAGAGGTACA---TGTTT

Phylogenetics

Darwin, 1837



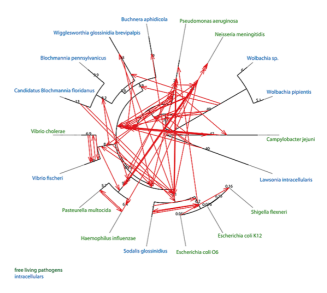
Haeckel, 1866



Phylogenetics

Darwin, 1837





Genome distance tree with lateral gene transfers

Information in (viral) molecular sequences

- Genetic distances among strains
- Phylogeny
 - → subtyping/classification
 - → identification of transmission clusters
 - ⇒ association with risk factors / traits
 - → forensics

- Dates of historical events
- Evolutionary processes
 - → recombination
 - → natural selection
- Epidemiological processes
 - → transmission rates
 - → movement among locations
- Phenotypic trait evolution?

HIV-1 (UK) ATC---TGCTAAAGCATATGACACAGAGGTACATAATGTTT
HIV-1 (USA) ATCGGATGCTAGAGCTTATGATACAGAGGTACA---TGTTT

Our goal

MOLECULAR SEQUENCES

Alignment Methods



BIOINFORMATICS

ALIGNMENT

Sequence Evolution Models Phylogenetic Methods



PHYLOGENETICS

EVOLUTIONARY TREE

(time scale = genetic distance)

Molecular Clock Models



PHYLOGENETICS

EVOLUTIONARY TREE

(time scale = years)

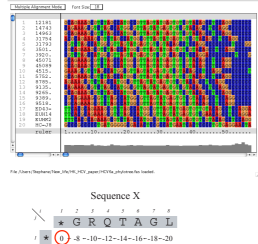
Phylodynamic Models

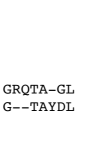


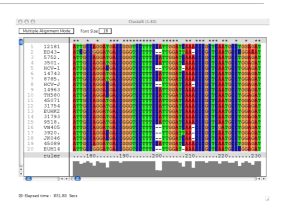
POPULATION GENETICS

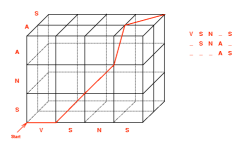
EPIDEMIOLOGY

Sequence alignment

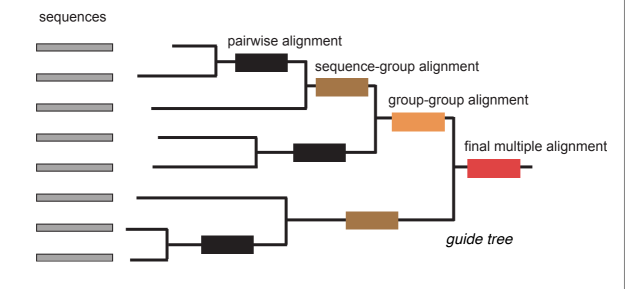








Progressive alignment



http://www.kuleuven.be/aidslab/phylogenybook/Table3.1.html

Genetic distances

```
ATGGGTGCGA GAGCGTCAGT TCTAACAGGG GGAAAATTAG ATCGCTGGGA ATGGGTGCGA GAGCGTCAGT ATTAAGCGGG GGAGAATTAG ATCGATGGGA
SIVcpz
HIV-1
                AAAAGTTCGG CTTAGGCCCG GGGGAAGAA AAGATATATG ATGAAACATT AAAAATTCGG TTAAGGCCAG GGGGAAAGAA AAAATATAAA TTAAAACATA
SIVcpz
HIV-1
                TAGTATGGGC AAGCAGGGAG CTGGAAAGAT TCGCATGTGA CCCCGGGCTA
TAGTATGGGC AAGCAGGGAG CTAGAACGAT TCGCAGTTAA TCCTGGCCTG
SIVcpz
HIV-1
                ATGGAAAGTA AGGAAGGATG TACTAAATTG TTACAACAAT TAGAGCCAGC
TTAGAAACAT CAGAAGGCTG TAGACAAATA CTGGGACAGC TACAACCATC
SIVcpz
HIV-1
                TCTCAAAACA GGCTCAGAAG GACTGCGGTC CTTGTTTAAC ACTCTGGCAGCCTTCAGACA GGATCAGAAG AACTTAGATC ATTATAAT ACAGTAGCAA
HIV-1
                 TACTGTGGTG CATACATAGT GACATCACTG TAGAAGACAC ACAGAAAGCT
SIVcpz
HIV-l
                 CCCTCTATTG TGTGCATCAA AGGATAGAGA TAAAAGACAC CAAGGAAGCT
                CTAGAACAGC TAAAGCGGCA TCATGGAGAA CAACAGAGCA AAACTGAAAG
TTAGACAAGA TAGAG-GAA ----GAGCA AAACAAAAGT AA---GAAAA
SIVcpz
HIV-1
                TAACTCAGGA AGCCGTGAAG GGGGAGCCAG TCAAGGCGCT AGTGCCTCTGAAGCACAGCA AGC---AG CAGCTGACA- -CAGGACAC- AG--CAGC--
SIVcpz
HIV-1
SIVcpz
                CTGGCATTAG TGGAAATTAC
HIV-1
                CAGG--TCAG CCAAAATTAC
```

chimpanzee SIV vs HIV-1 envelope gene

Not all mutations are equally likely

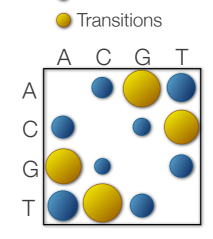
- some point substitutions are more likely to occur than others: transitions are more likely than transversions
 - transitions:

purine or

pyrimidine → pyrimidine **A**→**G C**→**T**

transversions:

 $A \leftrightarrow C$ $A \leftrightarrow T$ $G \leftrightarrow C$ $G \leftrightarrow T$

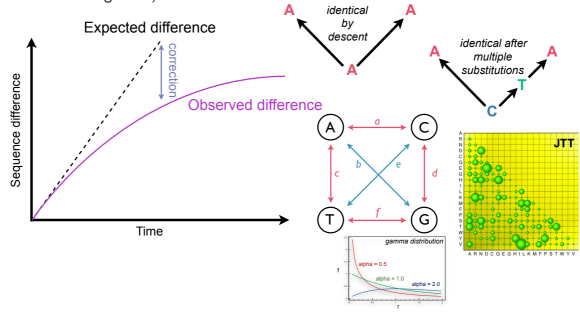


Transversions

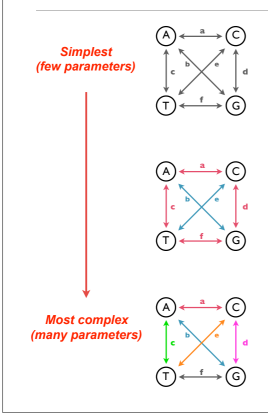
Unambiguous changes on most parsimonious tree of Ciliate SSUrDNA

Substitution models

 During evolution, 'multiple hits' can occur at a single position: the evolutionary distance is almost always larger than the dissimilarity (% nt or aa divergence)



Nucleotide substitution models



- 1. Base frequencies are equal and all substitutions are equally likely (Jukes-Cantor) (a=b=c=d=e=f)
- 2. Base frequencies are equal but transitions and transversions occur at different rates (Kimura 2-parameter)

(a=c=d=f, b=e)

3. Unequal base frequencies and transitions and transversions occur at different rates (Hasegawa-Kishino-Yano)

(a=c=d=f, b=e)

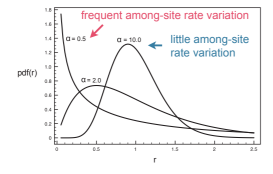
4. Unequal base frequencies and all substitution types occur at different rates (General Reversible Model)

(a, b, c, d, e, f)

Does this matter?

Estimated genetic distances between SIVcpz and HIVlai, under different substitution models:

Observed % mismatches = 0.406 JC (Jukes-Cantor) = 0.586 HKY (Hasegawa-Kishino-Yano) = 0.611 GTR (General Time Reversible) = 0.620 GTR + gamma = 1.017



Gene	α
Prolactin	1.37
Albumin	1.05
C-myc	0.47
Ctyochrome β (mtDNA)	0.44
Insulin	0.40
D-loop (mtDNA)	0.17
12S rRNA (mtDNA)	0.16

Phylogenetic reconstruction



Alignment Methods



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PHYLOGENETICS

EVOLUTIONARY TREE

(time scale = genetic distance)

Molecular Clock Models



EVOLUTIONARY TREE

(time scale = years)

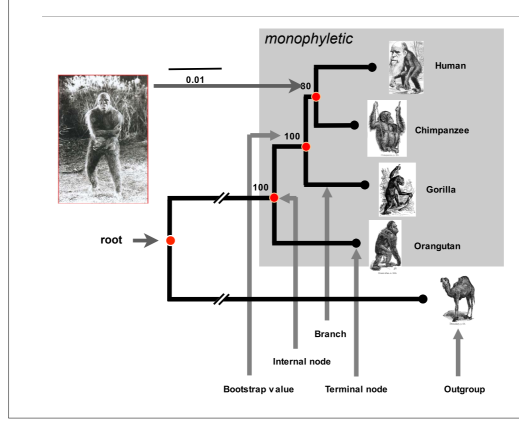
Phylodynamic Models

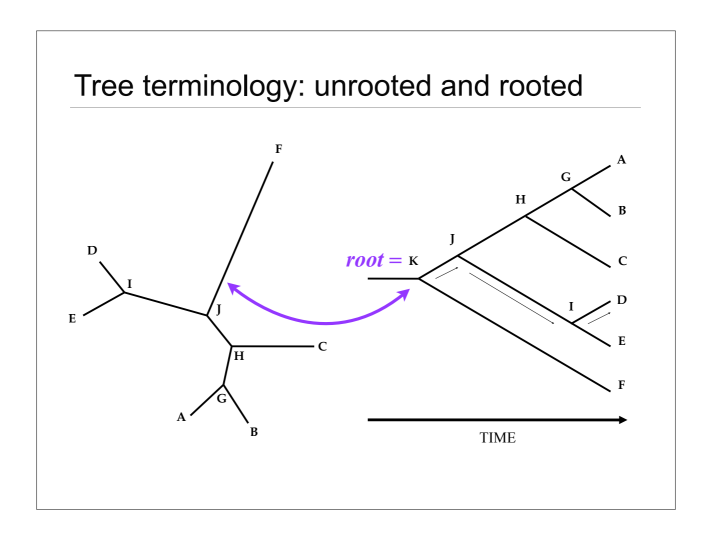


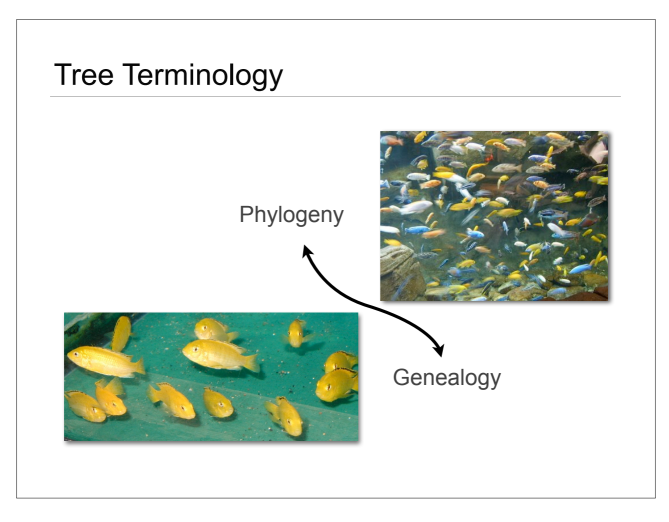
POPULATION GENETICS

EPIDEMIOLOGY

What is a tree?







Phylogenetic reconstruction

- **CLUSTERING APPROACHES:** These begin with a genetic distance between each pair of sequences. A 'clustering algorithm' then transforms the genetic distances into a tree.
 - e.g. UPGMA, Neighbour-Joining
 - · Simple, faster.
 - No measure of how good the estimated tree is (non-statistical)
- **OPTIMALITY METHODS**: These define a score for each possible tree. 'Search algorithms' are then used to find the tree with the highest score.
 - e.g. Parsimony, Maximum Likelihood (& Bayesian Inference)
 - More complex, slower. Search may not locate the 'best' tree.
 - Quality of each tree can be directly compared (statistical)

Phylogenetic reconstruction

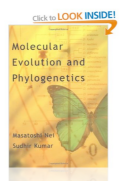
For n taxa, there are:

$$(2n-3)!/[(2^{n-2})*(n-2)!]$$
 rooted, binary trees

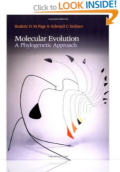
# taxa	# trees	
4 5 6	15 105 945	enumerable by hand enumerable by hand on a rainy day enumerable by computer
7 8	10395 135135	still searchable very quickly on computer a bit more than the number of hairs on your head
9 10	2027025 34459425	population of Glasgow ≈ upper limit for exhaustive searching; about the number of possible combinations of numbers in the National Lottery
20	8.20 × 10 ²¹	≈ upper limit for branch-and-bound searching
48 136	3.21 × 10 ⁷⁰ 2.11 × 10 ²⁶⁷	≈ the number of particles in the universe =number of trees to choose from in the "Out of Africa" data (Vigilant et al., 1991)

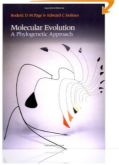
Phylogenetic inference: books





Click to LOOK INSIDE!





Click to LOOK INSIDE!

INFERRING PHYLOGENIES



- Yang Z. (2003). Computational Molecular Evolution. Oxford University Press
- Nei M & Kumar S. (2000). Molecular Evolution and Phylogenetics. Oxford University Press.
- Page RDM & Holmes EC. (1998). Molecular Evolution: A Phylogenetic Approach. Blackwell Science Ltd, Oxford.
- Yang Z (2014) Molecular Evolution: A Statistical Approach
- Bayesian Phylogenetics: Methods, Algorithms, and Applications. Chen M-H, Kuo L. and Lewis PO. Chapman & Hall/CRC.
- Lemey P, Salemi M & Vandamme A-M. (2009). The Phylogenetic Handbook, 2nd Edition. Cambridge University Press.
- Felsenstein J. (2003). Inferring phylogenies. Sinauer

Computer Software: http://evolution.genetics.washington.edu/phylip/software.html

PhylodynamicsTM **EPIDEMIC DYNAMICS** GENETIC DIVERSITY (mathematical epidemiology) (phylogenetics & molecular evolution) NATURAL SELECTION (population genetics & immunology)

Unifying principle

"Rapidly evolving pathogens are unique in that their ecological and evolutionary dynamics occur on the same timescale and can therefore potentially interact."

Pybus & Rambaut (2009) Nat. Rev. Genetics 10:540-50

Fundamental Phylodynamic Questions

- How genetically diverse is a pathogen population?
- How do pathogen genomes change through time?
- How does pathogen genetic diversity vary through time and space?
- What are the effects of pathogen genetic diversity on virulence, transmissibility, resistance to treatment, etc.

Specific questions

- When did a epidemic start?
- Where did it come from?
- How fast is it transmitting?
- In what direction is it spreading?
- Are hosts X, Y & Z epidemiologically linked?
- Of how many strains is the epidemic composed?
- Are strains associated with particular transmission routes?
- What adaptations has it accrued?

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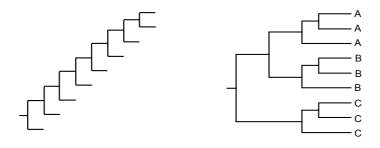
Measuring sequence diversity

- Not as straightforward as you might think...
- Are your pathogen sequences all sampled at the same time?

If sequences not sampled over time it's difficult to separate the effects of diversity and divergence on genetic diversity.

Are you measuring sample diversity or population diversity?

The former is simply a summary of your data, the latter is an inference about the population you have sampled. Sequences should be sampled randomly to estimate the latter.



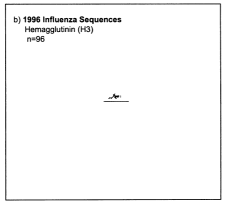
Measuring sequence diversity

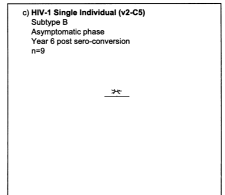
Are you studying an inter-host or intra-host population?
 For the former, each sequence represents a different infection.
 For the latter, each sequence represents a different virion within

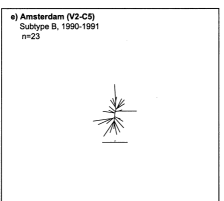
For the latter, each sequence represents a different virion within an infected individual. The measure of diversity must be interpreted accordingly.

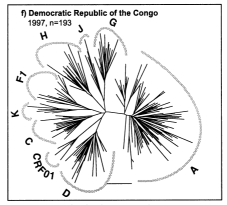
- How do we deal with intra-host diversity when studying the inter-host level?
- Intra-host diversity is low for most acute infections (e.g. influenza) but can be high for chronic infections (e.g. HIV).

Example: diversity of HIV-1 versus influenza



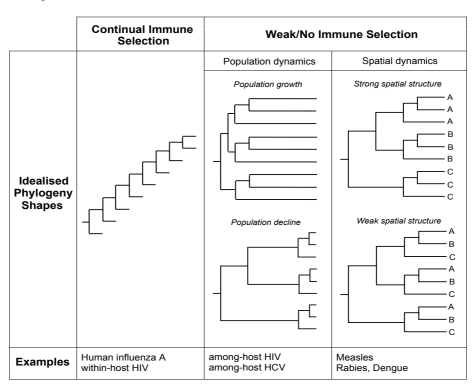






Scale bar represents a genetic distance of 0.1 substitutions per site. Korber et al. 2001. British Medical Bulletin 58:19-42

Phylodynamic Patterns

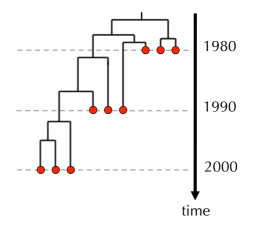


Fundamental Phylodynamic Questions

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'Phylodynamic' Data

- Pathogen genomes are sampled at different points in time and from different locations.
- Hence transmission history is estimated on a real time-scale (e.g. years).



MOLECULAR SEQUENCES





BIOINFORMATICS

ALIGNMENT

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EVOLUTIONARY TREE

(time scale = genetic distance)

Molecular Clock Models



PHYLOGENETICS

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(time scale = years)

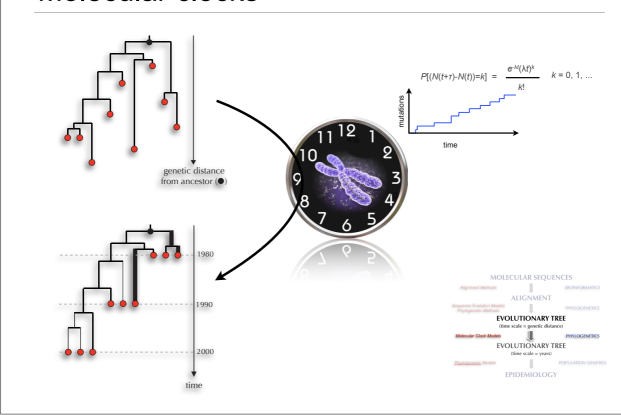
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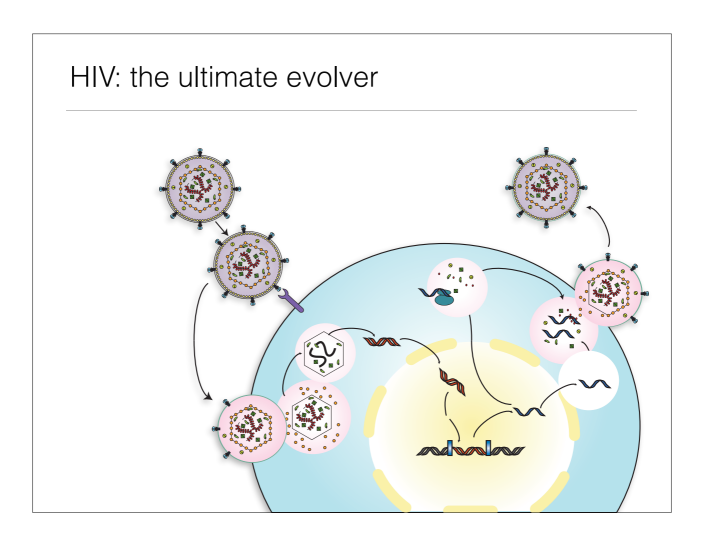


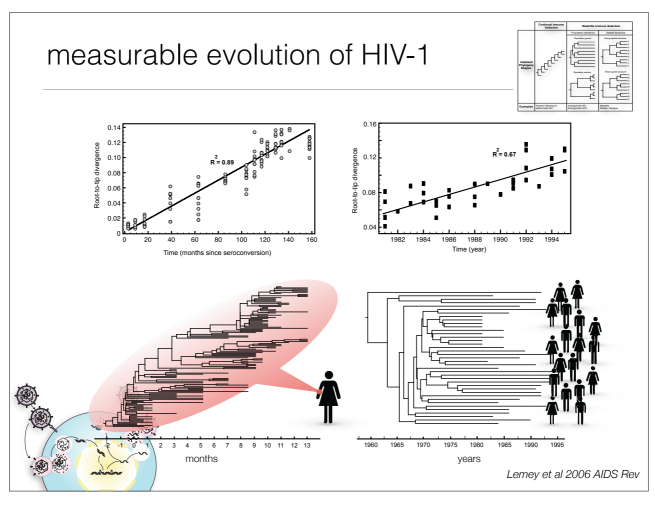
POPULATION GENETICS

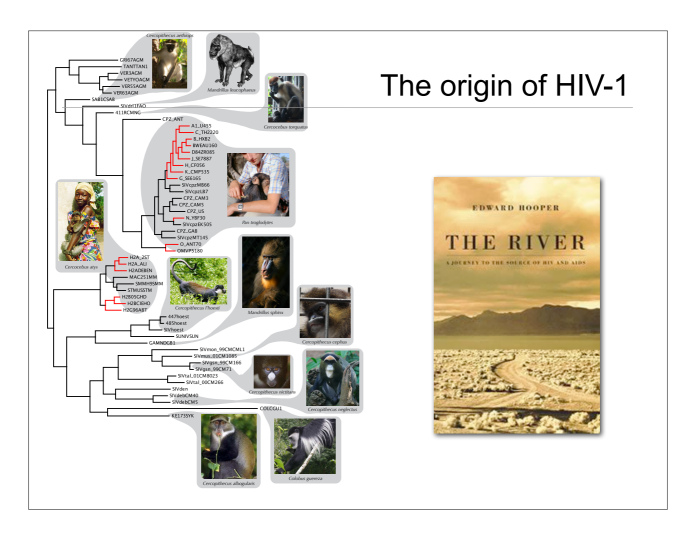
EPIDEMIOLOGY

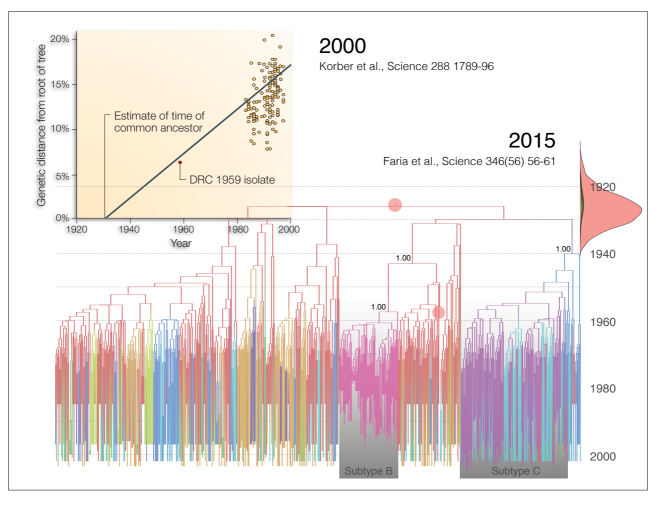
Molecular clocks





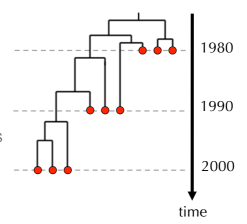


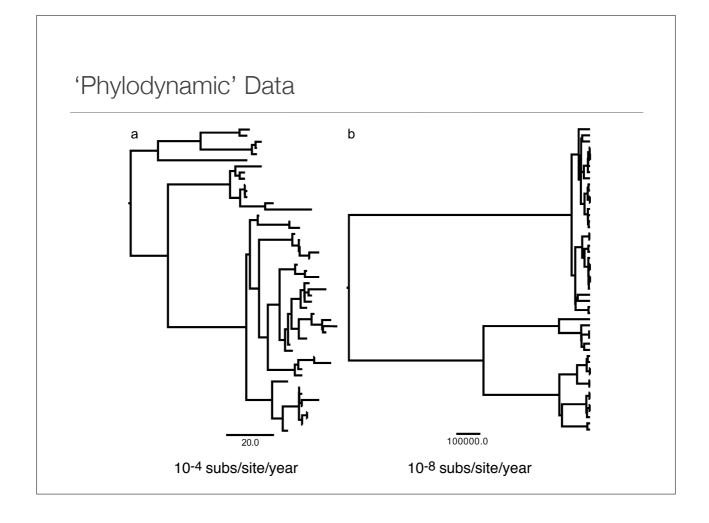


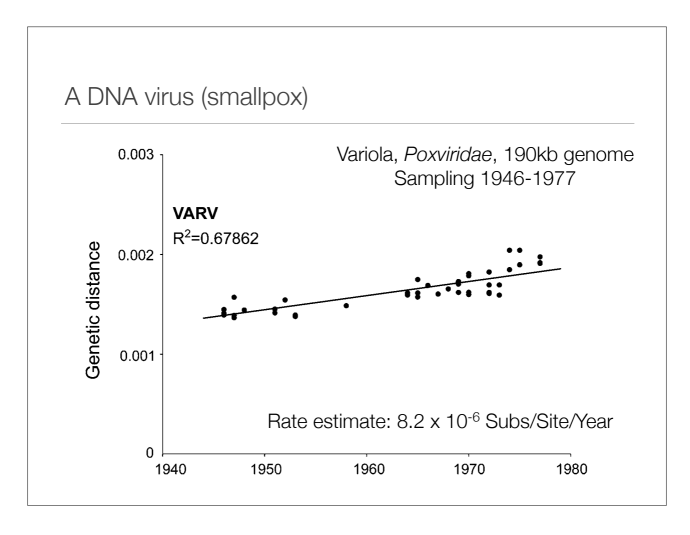


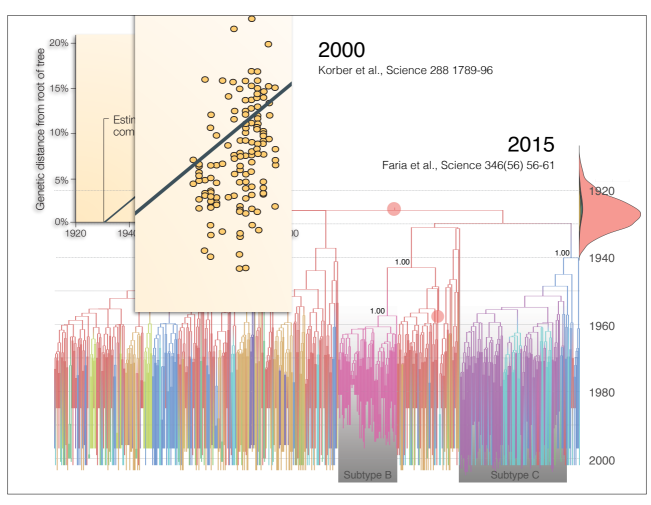
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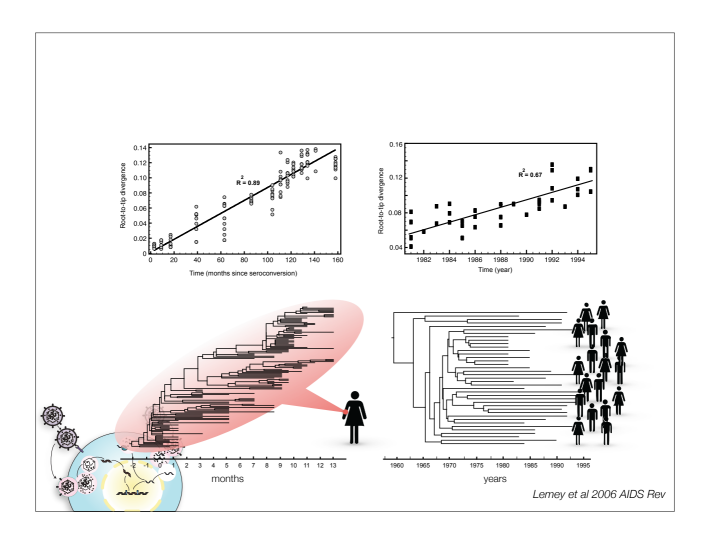
- Pathogen genomes are sampled at different points in time and from different locations.
- Hence transmission history is estimated on a real time-scale (e.g. years).
- The ability to genetically distinguish sequences sampled at different times depends on:
 - (i) the rate of evolution of the gene/ genome that is obtained
 - (ii) the length of time between samples
 - (iii) the sequence length of the gene/ genome that is obtained

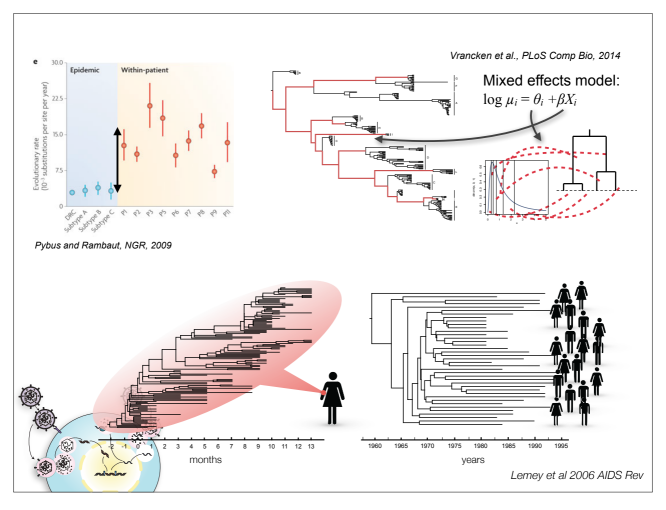


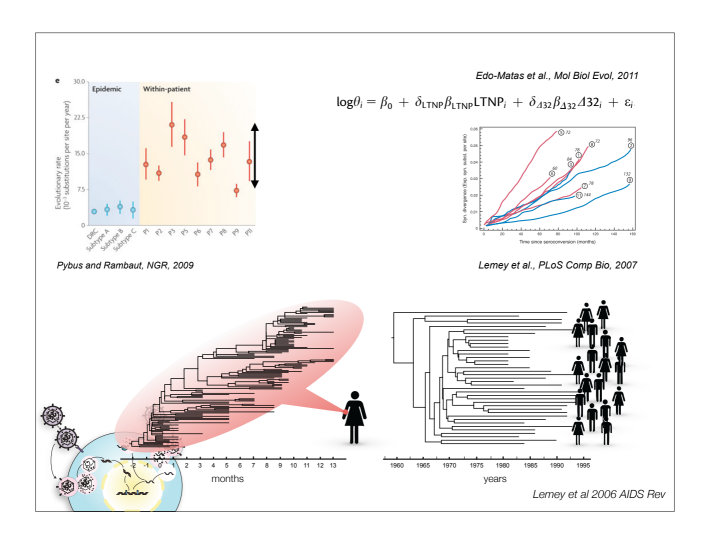


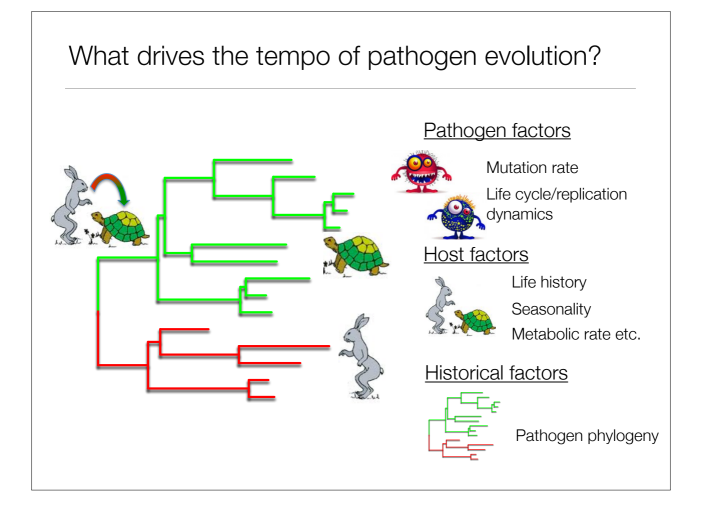






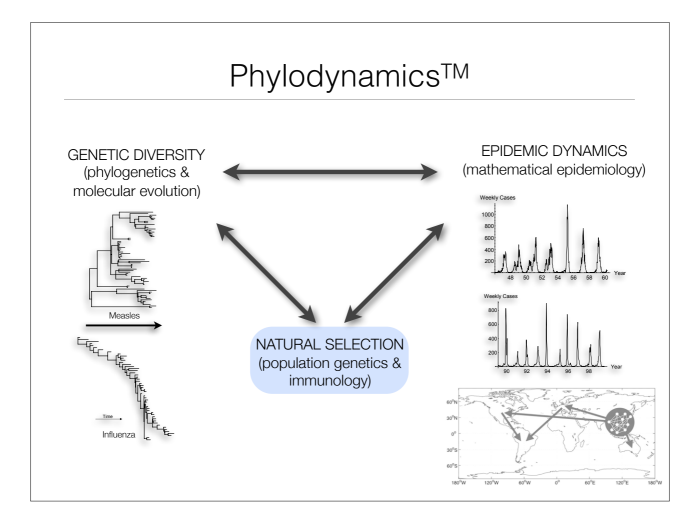






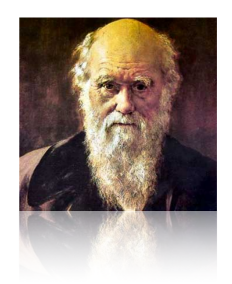
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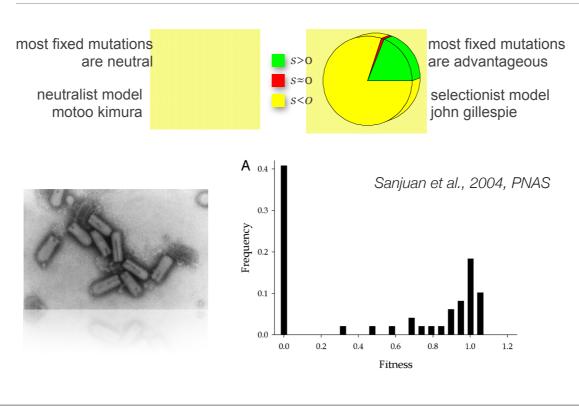


Evolutionary processes: natural selection

- "the preservation of favourable variations and the rejection of injurious variations, i call natural selection. variations neither useful nor injurious would not be affected by natural selection, and would be left a fluctuating element"
 - darwin, the origin of species

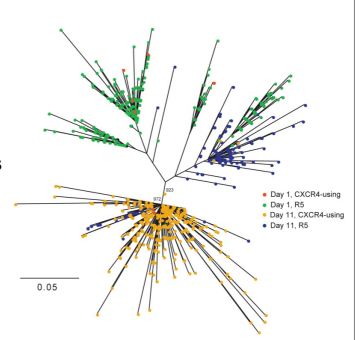


Evolutionary processes: natural selection



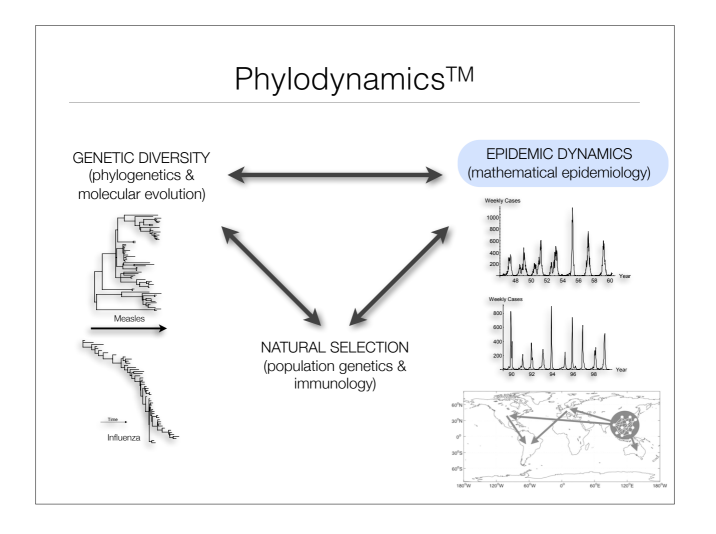
Evolutionary processes: natural selection

- Immune escape (antibodies*, T-cells*, innate immune responses)
- Antiviral drug resistance
- Vaccine escape mutations
- Cell & tissue tropism
- Inter-host viral transmission (i.e. for viral emergence)





module 15: Pathogen evolution, selection and immunology



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