# Coalescent Modeling for Distributions of Alleles

#### **Biostatistics 666**

# Last Lecture: Introduction to the Coalescent

- Coalescent approach
  - Proceed backwards through time.
  - Model the genealogy of sample of sequences.

# Infinite sites model

- All mutations distinguishable.
- No reverse mutation.

# Some key ideas ...

- Probability of coalescence events
- Length of genealogy and its branches
- Expected number of mutations
- Parameter θ which combines population size and mutation rate

# **Building Blocks...**

 Probability of sampling distinct ancestors for *n* sequences

$$P(n) = \prod_{i=1}^{n-1} \left( 1 - \frac{i}{N} \right) \approx 1 - \frac{\binom{n}{2}}{N}$$

 Coalescence time t is approximately exponentially distributed

# Some Key Results...

• Coalescence Time (population size units)  $E(T_j) = 1/\binom{j}{2}$ 

Total Tree Length (population size units)

$$E(T_{tot}) = \sum_{i=1}^{n-1} \frac{2}{i}$$

# Some More Key Results ...

Expected Number of Polymorphisms

For a diploid sample

$$E(S) = 4N\mu \sum_{i=1}^{n-1} 1/i = \theta \sum_{i=1}^{n-1} 1/i$$

For an haploid sample

$$E(S) = 2N\mu \sum_{i=1}^{n-1} 1/i = \theta \sum_{i=1}^{n-1} 1/i$$

# Estimating $\theta$

- Number of variants S can be used to estimate  $\theta$ 
  - Expected S is simply  $\theta E(T_{tot})$
  - To estimate θ, divide by S expected length of genealogy

$$\hat{\theta} = \frac{S}{\sum_{i=1}^{n-1} 1/i}$$

- Could then be used to:
  - Estimate N, if mutation rate  $\mu$  is known
  - Estimate μ, if population size N is known

# Alternative Estimator for $\theta$ ...

 Count pairwise differences between sequences

Compute average number of differences

$$\widetilde{\theta} = \binom{n}{2}^{-1} \sum_{i=1}^{n} \sum_{j=i+1}^{n} S_{ij}$$

# Var( $\hat{\boldsymbol{\theta}}$ ) as a function of *n*



# Var( $\hat{\boldsymbol{\theta}}$ ) as a function of *n*



# Today ...

#### More applications of the coalescent

# Predicting allele frequency distributions Using simulations

Using simulations

# Modeling the distribution of S

Using analytical calculations

# A Coalescent Simulation ...

Let's consider tracing the ancestry of 4 sequences

#### When n = 4

Probability of Coalescent Event

$$P(4) \approx \binom{4}{2} / 2N$$

Time to Next Coalescent Event

$$T(4) \approx 2N / \binom{4}{2}$$

Sample time from exponential distribution

Pick two sequences at random to coalesce

#### Next n = 3 ...

Let's assume that sequences 3 and 4 are selected ...

Then, we repeat the process for a sample of 3 sequences



#### Next n = 2 ...

Let's assume that sequences 1 and 2 are selected to coalesce

Then, we repeat the process for a sample of 2 sequences



# **The Simulated Coalescent**



# A Coalescent Simulation ...



# A Coalescent Simulation ...



# **Frequency Spectrum**

 Repeating the simulation multiple times, would give us a predicted mutation spectrum.



# Frequency Spectrum (n = 10)



# Frequency Spectrum (n = 100)



# **Frequency Spectrum**

- Constant size population
- Exponentially growing population

#### Most variants are rare

- For n = 100, ~44% of variants occur < 5/100.</p>
- For n = 10, ~35% of variants observed once.

# **Mutation Spectrum**

#### Depends on genealogy

- Population Size
- Population Growth
- Population Subdivision
- Does not depend on
  - Mutation rate!

# **Deviations from Neutral Spectrum**

- When would you expect deviations from the spectra we described?
- What would you expect for ...
  - A rapidly growing population?
  - A population whose size is decreasing?
- Why?

# **Effect of Polymorphism Type**



# **Frequency Spectrum of Protein Altering Variants**



Exome Chip Consortium (2011)

# **Number of Mutations**

- Can be derived from coalescent tree
  - What are the key features?
- Analytical results possible
  - Trace back in time until MRCA, tracking mutation events

# **Sample of Two Sequences**

- Track coalescences and mutations
  - Probability of a coalescent event?
    - Depends on population size …
  - Probability of a mutation?
    - Depends on mutation rate ...
- Proceed backwards until either occurs...
  - Conditional probability for each outcome?

# **Two Identical Sequences**

$$P_2(S \text{ is } 0) \approx \frac{P_{CA}}{P_{CA} + P_{mut}}$$
$$= \frac{1/2N}{1/2N + 2\mu}$$
$$= \frac{1}{1+\theta}$$

# Full distribution of S...

Probability that first *j* events are mutations...

# $P_2(j) = \left(\frac{\theta}{1+\theta}\right)^j \left(\frac{1}{1+\theta}\right)^j$

#### Example...

- 2 sequences
- Population size N = 25,000
- Mutation rate  $\mu = 10^{-5}$

# Probability of 0, 1, 2, 3... mutations

# And for multiple sequences...

 Describe number of mutations until the next coalescence event

- Proceed back in time, until:
  - One of *n* sequences mutates...
  - A coalescent event occurs...
    - Then track mutations in (n-1) sequences

#### Formulae ...



#### Example...

- 3 sequences
- Population size N = 25,000
- Mutation rate  $\mu = 10^{-5}$

## Probability of 0, 1, 2, 3... mutations

#### **Number of Mutations**



# So far ...

# One homogeneous population

- Coalescence times
- Number of mutations
  - Expectation
  - Distribution
- Spectrum of mutations
- Several assumptions, including ...
  - No recombination

# **Next: Models w/ Recombination**

- No recombination
  - Single genealogy
- Free recombination
  - Two independent genealogies
  - Same population history
- Intermediate case
  - Correlated genealogies

# **Recommended Reading**

## Richard R. Hudson (1990)

Gene genealogies and the coalescent process

Oxford Surveys in Evolutionary Biology, Vol. 7. D. Futuyma and J. Antonovics (Eds). Oxford University Press, New York.