## Summer Institute in Statistical Genetics

 University of Queensland, Brisbane Module 5: Population Genetic Data Analysis
## The Coalescent Model

David Balding<br>Professor of Statistical Genetics University of Melbourne, and University College London

Feb 9, 2017

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## Going backwards in time: coalescent models

DNA sequences at the same locus from different individuals are dependent:

- differing amounts of common ancestry;
- so differing levels of correlation among the sequences.

Valid inferences from DNA sequence data, e.g. about mutation rates or about the location of a gene of interest, may require modelling the relationships among the sequences.

- Incorrectly assuming independence can lead to understatement of variances of estimators - the effect can be large.
- Sometimes the relationships among the sequences are crucial e.g. for inferences about population histories.

A natural way to describe both the pattern of shared ancestry and the resulting correlations is via a genealogical tree (similar to a phylogenetic tree but for genes within a population, rather than from different species).

## Coalescent models

Possible genealogy of a sample of 6 homologous sequences, showing two mutation events. The time arrow points backwards: e.g. $t_{6}$ denotes the most recent coalescent event, when the number of lineages decreased (going back in time) from 6 to 5 .


The (standard) coalescent is a model for the genealogy underlying a sample of $n$ genes at a neutral, non-recombining locus drawn from a large, random-mating, constant-size population.

- Leaves of the tree $\Leftrightarrow$ observed DNA sequences;
- going up the tree $\Leftrightarrow$ tracing the ancestry of the sequences;
- branches merge, or "coalesce", when the descendant sequences first share a common ancestor;
- the root of the tree corresponds to the Most Recent Common Ancestor (MRCA) of all the sequences in the sample.
Under the coalescent model, the time during which the tree has $j$ distinct branches has the exponential distribution with parameter $j(j-1) / 2$ (we write $\operatorname{Exp}(j(j-1) / 2)$, NB mean $=1 /$ parameter). The times for different $j$ are independent. Here, one unit of "coalescent" time corresponds to $N G / \sigma^{2}$ years, where $N=$ (effective) population size,
$G=$ generation time,
$\sigma^{2}=$ variance in number of offspring (below assume $\sigma^{2}=1$ ).


Let $T_{n}$ and $L_{n}$ denote the height and the total branch length of a coalescent tree with $n$ leaves. Then

$$
\begin{aligned}
& E\left[T_{n}\right]=2(1-1 / n) \quad E\left[L_{n}\right]=\sum_{j=1}^{n-1} \frac{2}{j} \approx 1+2 \log (n) \\
& \operatorname{Var}\left[T_{n}\right]=\sum_{j=2}^{n} \frac{8}{j^{2}}-4\left(\frac{n-1}{n}\right)^{2} \quad \operatorname{Var}\left[L_{n}\right]=\sum_{j=1}^{n-1} \frac{4}{j^{2}}
\end{aligned}
$$

Features of the coalescent model:

- the mean time in which the sample has exactly two ancestors is more than half $\mathrm{E}\left[T_{n}\right]$, the mean total time since the MRCA (this can lead to bimodality in datasets);
- the variance of $T_{n}$ is large relative to its mean; the largest contribution to $\operatorname{Var}\left[T_{n}\right]$ arises from the interval in which the sample has just two ancestors;
- $E\left[L_{n}\right]$, the mean total branch length of tree (which is roughly the total amount of independent information in the data) grows only like $\log (n)$, not $n$ as would be the case for a random sample.
- Although $\mathrm{E}\left[L_{n}\right]$ continues to increase with $n, \operatorname{Var}\left[L_{n}\right]$ does not.

These observations have big implications for patterns of DNA sequence variation along the genome.


## Coalescent with mutation

Mutations occur along the branches of a coalescent tree uniformly at random with rate $\theta / 2$, where $\theta=2 N \mu$ and $\mu$ is the mutation rate per sequence per generation. Given $L_{n}$, the total branch length of the tree, the number $S_{n}$ of mutations has the Poisson distribution with mean $\theta L_{n} / 2$. The unconditional expectation is

$$
E\left[S_{n}\right]=\frac{\theta}{2} E\left[L_{n}\right]=\theta \sum_{j=1}^{n-1} \frac{1}{j}
$$

If $\mu$ is small, it may be reasonable to assume the infinite sites model: every mutation is at a distinct site. Then $S_{n}$ is just the number of variable sites and a natural estimator of $\theta$ is Watterson's estimator $\hat{\theta}_{W}=S_{n} / \sum_{j=1}^{n-1} \frac{1}{j}$, which is unbiased but the variance decreases like $1 / \log (n)$, much slower than $1 / n$ for estimators obtained from random samples.

- Even very big samples don't give very accurate estimates.

The standard coalescent model

The coalescent with population growth

## Statistical inference under the coalescent

## Simulation-based inference under the coalescent

Example: the Irish giant and AIP acromegaly mutation

- Extensions of the standard model can allow for changes in population size, population subdivision, natural selection and recombination.
- The simplest extension is to incorporate population growth.
- Suppose that the population size $N t$ generations ago was $N \lambda(t)$, where $N$ denotes the current effective population size, so that $\lambda(0)=1$. Scaling time by $N$ as for the standard coalescent, the waiting time for the $j$ th coalescence event is now given by

$$
\begin{equation*}
P\left(w_{j}>t\right)=\exp \left(-\frac{j(j-1)}{2} \Lambda(t)\right) \tag{1}
\end{equation*}
$$

where $\Lambda(t)=\int_{0}^{t} \mathrm{~d} s / \lambda(s)$.

- When the population size is large (i.e. $\lambda(t)$ is large), $\Lambda(t)$ increases only slowly with $t$, corresponding to the fact that coalescences rarely occur.


## Four realisations of the coalescent model with mutation;

 sample size $n=6$; exponential growth, $R=100$.

The time scaling has been chosen so that $\mathrm{E}\left[L_{n}\right]$ is approximately the same as for the standard coalescent.

The standard coalescent arises in the case that $\lambda(t) \equiv 1$ and $\Lambda(t)=t$. Exponential growth/decline forward in time at rate $r$ per generation corresponds to

$$
\begin{equation*}
\lambda(t)=\exp (-R t) \quad \Lambda(t)=\frac{\exp (R t)-1}{R} \tag{2}
\end{equation*}
$$

where $R=N r$.

- Large $R$ implies rapid growth forward in time $\Leftrightarrow$ rapid decline backward in time: relatively few coalescences occur in the recent past because the population size is large.
- In the limit as $R \uparrow \infty$ we obtain a "star genealogy": all coalescence events occur at about the same time and hence observed haplotypes are independent given the ancestral haplotype.

The following plots show histograms of the numbers of mutations under 10000 realisations of (a) standard coalescent model and (b) coalescent with growth, in each case compared with expected values under the Poisson distribution with matching mean.

## Distribution of mutation count: standard coalescent



## Distribution of mutation count: coalescent with growth



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Example: the Irish giant and AIP acromegaly mutation

- Coalescent models without mutation specify a prior distribution for the genealogy underlying a set of DNA sequences, given the sample sizes but not the sequence data.
- Coalescent models with mutation can give predictive distributions for properties of sequence data expected under different models (with growth, structure, selection etc).
- However, often what we want to do is to infer properties of the underlying model (such as the mutation rate or time since most recent common ancestor (TMRCA)) given observed data.
One way to proceed is to seek the posterior distribution of parameters of interest, given the observed data, the coalescent model as prior for the genealogical tree, and assumed prior distributions for evolutionary parameters.
- To obtain the posterior from the prior, use Bayes Theorem:

$$
\operatorname{Pr}(\theta \mid D)=\frac{\operatorname{Pr}(D \mid \theta) \operatorname{Pr}(\theta)}{\operatorname{Pr}(D)}
$$

where $\operatorname{Pr}(D)=\int \operatorname{Pr}(D \mid \theta) \operatorname{Pr}(\theta) d \theta$.

## Exact Inference for TMRCA when $n=2$

- We assume the standard coalescent model with infinite-sites mutation and suppose that $\theta$ is known;
- the unknown of interest is the coalescence time, or TMRCA of the two sequences, let's call it $t_{2}$.
Given $t_{2}=t$, the number of segregating sites $S$ has a Poisson distribution with parameter $\theta t$ :

$$
\begin{equation*}
\operatorname{Pr}\left(S=s \mid t_{2}=t\right)=\frac{1}{s!}(\theta t)^{s} \exp (-\theta t) \tag{3}
\end{equation*}
$$

By Bayes theorem we obtain the posterior pdf of $t_{2}$ :

$$
\begin{equation*}
\mathrm{p}\left(t_{2}=t \mid S=s\right)=C(\theta t)^{s} \exp (-(1+\theta) t) \tag{4}
\end{equation*}
$$

where $C$ is a constant (does not depend on $t$ ). The RHS of (4) has the form of the Gamma $(1+s, 1+\theta)$ probability density function, and it follows that (Tajima, 1983):

$$
\begin{equation*}
\mathrm{E}\left[t_{2} \mid S=s\right]=\frac{1+s}{1+\theta} \quad \text { and } \quad \operatorname{Var}\left[t_{2} \mid S=s\right]=\frac{1+s}{(1+\theta)^{2}} \tag{5}
\end{equation*}
$$

which may be compared with prior moments $\mathrm{E}\left[t_{2}\right]=\operatorname{Var}\left[t_{2}\right]=1$.

- Noting that $\mathrm{E}[S]=\theta$, we see that if $s<\mathrm{E}[S]$ then $\mathrm{E}\left[t_{2} \mid S=s\right]<\mathrm{E}\left[t_{2}\right]$, and vice-versa.
- Data usually decreases the variance: $\operatorname{Var}\left[t_{2} \mid S=s\right]<\operatorname{Var}\left[t_{2}\right]$ unless $s$ is very large $\left(\geq 2 \mathrm{E}[S]+\mathrm{E}[S]^{2}\right)$.

The prior density curve and posterior curves for several values of $s$ when $\theta=1$ are shown on next slide.


Density curves for TMRCA ( $t_{2}$, in coalescent units) when $n=2$, $\theta=1$. Prior: $\operatorname{Gamma}(1,1) \equiv \operatorname{Exp}(1)$; posteriors: $\operatorname{Gamma}(1+s, 2)$.

## Comparison with classical estimator

- A natural estimator of $t_{2}$ within the framework of classical statistics is the method-of-moments estimator

$$
\hat{t}_{2}=S / \theta
$$

for which the mean square error (MSE) is

$$
\operatorname{MSE}(S / \theta)=\mathrm{E}_{t_{2}}\left[\mathrm{E}_{S \mid t_{2}}\left[\left(S / \theta-t_{2}\right)^{2}\right]\right]=1 / \theta
$$

- Uniformly larger than $1 /(1+\theta)$, the MSE of $\mathrm{E}\left[t_{2} \mid S\right]$.

The use of prior distributions in statistical inference has been controversial, but here the prior is based on solid ground: the coalescent model that has been shown to provide a good approximation in many real populations.

- Additional information from prior $\Rightarrow$ more precise inferences. An additional advantage of the Bayesian paradigm for statistical inference is that we obtain a full posterior distribution which summarises all available information about the unknown TMRCA, rather than just a point estimator and its standard error.


## Exact inference for TMRCA when $S=0$

Dorit et al. (1995) sequenced a 729-bp fragment in $n=38$ human Y-chromosomes, observed $S=0$ and reported a TMRCA estimate of $\hat{t_{38}}=270 \mathrm{~K}$ years before present.

- One of the first explicit uses of coalescent theory for population genetics inference.
- A breakthrough! But unfortunately they made mistakes.

Donnelly et al. Science 1996, reply to Dorit:

- Assume no mutations in the underlying genealogy, then

$$
\begin{equation*}
\operatorname{Pr}\left(S=0 \mid \theta, w_{38}, w_{37}, \ldots, w_{2}\right)=\prod_{j=2}^{38} \exp \left(-j w_{j} \theta / 2\right) \tag{6}
\end{equation*}
$$

$w_{j}=$ length of time that the coalescent has exactly $j$ branches.

- The $w_{j}$ have independent $\operatorname{Exp}(j(j-1) / 2)$ prior distributions; from (6), posteriors are independent $\operatorname{Exp}(j(j-1+\theta) / 2)$.


Fig. 1. Summary statistics for the conditional distribution, under the coalescent model, of the time $T$ (in years) since the common ancestor, given a sample of 38 sequences which exhibit no variabil$i t y$, as a function of $N$, the effective population size. The generation time is assumed to be 20 years, and the mutation rate of the sequenced region per generation is taken to be $1.96 \times 10^{-5}$. Conditional distribution of $T$ follows from equation 5.2 in (7).

Thus posterior mean and variance of $t_{38}=\sum_{j=2}^{38} w_{j}$ are:
$\mathrm{E}\left[t_{38} \mid \theta, S=0\right]=$ $\sum_{j=2}^{38} \frac{2}{j(j-1+\theta)}$ $\operatorname{Var}\left[t_{38} \mid \theta, S=0\right]=$ $\sum_{j=2}^{38} \frac{4}{j^{2}(j-1+\theta)^{2}}$.

The observation of $S=0$ reduced the mean of $t_{38}$ by only around $20 \%$ to $40 \%$ from its prior mean for plausible $N$ values.

## Inference about $\theta, N$ and $\mu$

- Integrating over the $w_{j}$ in (6) we obtain the likelihood for $\theta$ :

$$
\begin{equation*}
\operatorname{Pr}(S=0 \mid \theta)=\prod_{j=2}^{n} \frac{j-1}{j-1+\theta} \tag{7}
\end{equation*}
$$

The MLE is $\hat{\theta}=0$, which is non-sensical a priori.

- This defect of the MLE can be avoided by reporting a posterior $95 \%$ highest-density interval, using either an improper uniform prior for $\theta$ or a proper, informative prior.
- An additional advantage to a Bayesian approach is that it becomes possible to report inferences about $N$ and $\mu$ separately (recall $\theta=2 N \mu$ ).
- The likelihood (7) only depends on $N$ and $\mu$ through their product. So the data do not help distinguish them, but an informative prior distribution, if available, can.


## Donnelly et al. Science 1996: reply to Dorit



- Here, $\mu=1.96 \times 10^{-5}$ and the generation time $G=20$ years.
- Modal values of $N$ are around 120 K years, but variance is wide.
- Note that inferences about $N$ and $\mu$ are always sensitive to the prior assumptions, whereas in the presence of sufficient data inferences about $\theta$ will be robust to the prior.

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Example: the Irish giant and AIP acromegaly mutation

## Rejection sampling

In most cases of interest exact inference under the coalescent is infeasible, but there are approximate methods based on simulation. A general approach to inference about $\theta$ given a sample of $n$ DNA sequences is as follows:

1. Simulate a coalescent tree with $n$ leaves,
2. simulate $\theta$ under an appropriate prior model,
3. simulate mutations along the branches of the tree according to a mutation model.
4. If the $n$ resulting sequences are sufficiently close to the observed sequences, accept the simulated $\theta$, otherwise reject.
5. The set of accepted $\theta$ values is approximately a sample from the posterior distribution of $\theta$ given the sequence data.
This is the core of the Approximate Bayesian Computation (ABC) method that has revolutionised population genetics over the past 15 years, allowing approximate inference under sophisticated models e.g. for population growth and structure.

- A key problem is to define "sufficiently close".
- In some settings the number of segregating sites $S$ captures most of the information in the sequence data.
- Conditional on $L$, the total branch length of the coalescent tree, $S$ has approximately a Poisson distribution with mean $\theta L / 2$.
- Therefore, the accept/reject step can be performed more efficiently using Poisson probabilities, without simulating a value of $S$ (Tavaré et al. 1997, Genetics).
- Coalescent tree and $\theta$ still need to be simulated unless $S=0$.


Figure 2. -Pre- and post-data density curves for $T_{10}$ with $\theta$ $=1 .-$, pre-data density; $\cdots, S_{10}=1 ;--, S_{10}=3 ;---, S_{10}$ $=5$.
qrej $=$ function(nacc=10000,nblk=round(nacc/2),nsamp=6,nsit=5,s=3) \# rejection inference about $N$, mu and TMRCA given
\# s segregating sites in nsamp sequences of length nsit
$\{$
count = 0
ns1 = nsamp-1
rate $=(n s 1: 1) *(n s a m p: 2) / 2$
acc $=$ matrix $(0,1,3)$
while (nrow (acc) <nacc+1)
\{
count $=$ count + nblk
$\mathrm{w}=\operatorname{matrix}(\operatorname{rexp}(\mathrm{ns} 1 * \mathrm{nblk}, \mathrm{rate}), \mathrm{ns} 1, \mathrm{nblk})$
TMRCA $=\operatorname{apply}(\mathrm{w}, 2$, sum $)$
$\mathrm{L}=\operatorname{apply}\left(\left(\right.\right.$ nsamp:2)$*_{\mathrm{w}}, 2$, sum $)$
u = runif (nblk)
$\mathrm{N}=\mathrm{rgamma}\left(\mathrm{nblk}, 5,10^{n}-3\right)$
$\mathrm{mu}=\operatorname{rgamma}\left(\mathrm{nblk}, 2,2 * 10^{\wedge} 4\right)$
ind $=u<\operatorname{dbinom}(s, n s i t, 1-\exp (-L * N * m u / n s i t)) / d b i n o m(s, n s i t, s / n s i t)$ acc $=\operatorname{rbind}(a c c, \operatorname{matrix}(c(N[i n d], m u[i n d], T M R C A[i n d]), 3))$
\}
list (count, acc $[-1$,$] )$
\}

## Rejection-sampling inference: exercise

Use the R code qrej (for quick rejection sampling) to perform inferences under the coalescent when $S$ segregating sites (default $=3$ ) are observed in $n$ sequences (default $=6$ ). qrej assumes:

- gamma $\left(5,10^{-3}\right)$ prior for $N$ (mean $\left.=5000, \mathrm{SD}=2236\right)$; and
- gamma( $2,2 \times 10^{4}$ ) prior for $\mu$ (mean $10^{-4}, \mathrm{SD}=7.07 \times 10^{-5}$ ). qrej returns a list of length 2 :

1. the number of iterations (must exceed nacc);
2. a matrix with 3 cols: accepted values of $N, \mu$ and TMRCA. To obtain a density plot e.g. for $N$, you can do:
plot(density(res[[2]][,1]),fro=0,to=15000)
and you should add an xlab to label the $x$-axis. You can also add a prior density:

$$
\text { lines (x,dgamma }(x, 5,0.001), \operatorname{lty}=2)
$$

where x is a vector of grid points between 0 and 15,000 .

- The plot shows some results from inference using qrej with default settings.
- Dashed curves: prior,
- Solid curves: posterior.
- You should try to replicate these plots and obtain a similar plot for TMRCA.


Density curves for mu



Density curves for theta


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## Radian et al. Human Mutation (2016). Coalescent-based estimate of number of carriers of AIP risk allele in Ireland.

AIP mutations cause autosomal dominant familial isolated pituitary adenomas (FIPA), most commonly manifesting as acromegaly or gigantism. Due to incomplete penetrance, the disease can also manifest as apparently sporadic pituitary adenoma (PA).

- Chahal et al. NEJM (2011): 5 carriers identified in mid-Ulster, including a proband case: a C18 "Irish giant".
- Coalescent simulation-based analysis predicted a large number of carriers concentrated in mid-Ulster
Population screening in mid-Ulster for AIP mutations:
- 81 carriers ( 30 affected, 18 pedigrees) identified in mid-Ulster.
- Low prevalence in Belfast ( $n=1000$ ), no carriers found in Republic of Ireland ( $n=2000$ ).
- Haplotype conservation suggested a recent TMRCA.

Now we seek to update predictions of the TMRCA and consequently the number of carriers not yet identified.


FIPA 6
FIPA 7
FIPA 8


Sp 7


|  | FIPA UK |  | FIPA <br> Romania |  | FIPA US (Italian) |  | Sp India |  | Sp Mexico |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | ${ }_{194}{ }_{19}$ |  |  |  | ${ }^{80}$ |  |
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| D115913 |  | 97 |  |  |  |  | 101 | 101 |  |  |
| ${ }^{\text {D } 11151249}$ | ${ }^{108}$ | ${ }_{118}^{126}$ |  | 116 128 | 124 | 108 128 | ${ }_{12}^{12}$ | 114 124 |  |  |
|  |  |  |  |  |  |  |  |  | 120 <br> 101 <br> 109 |  |
| Chrl1-67-7T-107 D11s4178 | $\underset{\substack{103 \\ 288}}{\substack{18 \\ \hline}}$ | $\underset{\substack{103 \\ 288}}{\substack{18}}$ | $c1002is$ | $\substack { 100 \\ \begin{subarray}{c}{120{ 1 0 0 \\ \begin{subarray} { c } { 1 2 0 } } \\{250} \end{subarray}$ |  | ${ }_{\text {c }}^{\substack{108 \\ 285}}$ | $c101265285$ |  |  | ${ }_{23}^{23}$ |
|  |  |  |  |  |  |  |  | ${ }^{\text {a }}$ |  |  |

Haplotypes on chromosome 11q12.2-13.3 of individuals carrying AIP R304*. Dark shading: haploblock shared by all lrish pedigrees; light shading: additional shared haploblocks. AIP alleles (black = wild-type, yellow $=$ R304*).

## Radian et al. (2016): Methods

- Haplotypes inferred from genotypes using PHASE.
- We performed exact coalescent inference for the fully conserved haplotype, the result of which became a prior for $A B C$ inference of the varyingly-shared haplotypes.
- Since we were concerned only with conserved haplotypes, recombination and mutation have the same effect (of destroying conservation) and so we treated recombination events like mutations and combined the two rates.
- The statistic used to compare simulated and observed datasets was the number of haplotypes sharing each genome segment (defined by consecutive short tandem repeat markers).
Results:
- TMRCA estimated at $2550(1275-5000)$ years.
- Forward simulations using TMRCA distribution predicted 432 ( $90-5175$ ) current carriers, including 86 affected assuming $20 \%$ penetrance.

