Coalescent-based association mapping and fine mapping of complex trait loci

Sebastian Zöllner

and

Jonathan K. Pritchard

Department of Human Genetics
The University of Chicago

October 7, 2004

1Address for correspondence: Department of Human Genetics, University of Chicago, 920 E 58th St–CLSC 507, Chicago IL 60637, USA; Email szoellne@genetics.uchicago.edu; phone 773 834-5243; fax 773 834-0505
Abstract

We outline a general coalescent framework for using genotype data in linkage disequilibrium-based mapping studies. Our approach unifies two main goals of gene mapping that have generally been treated separately in the past: detecting association (i.e., significance testing) and estimating the location of the causative variation. In order to tackle the problem, we separate the inference into two stages. First, we use Markov chain Monte Carlo to sample from the posterior distribution of coalescent genealogies of all the sampled chromosomes without regard to phenotype. Then, averaging across genealogies, we estimate the likelihood of the phenotype data under various models for mutation and penetrance at an unobserved disease locus. The essential signal that these models look for is that in the presence of disease susceptibility variants in a region, there is non-random clustering of the chromosomes on the tree according to phenotype. The extent of non-random clustering is captured by the likelihood, and can be used to construct significance tests or Bayesian posterior distributions for location. A novelty of our framework is that it can naturally accommodate quantitative data. We describe applications of the method to simulated data, and to data from a Mendelian locus (CFTR, responsible for cystic fibrosis) and from a proposed complex trait locus (Calpain-10, implicated in Type 2 diabetes).

Keywords: coalescent, fine mapping, complex disease, complex traits, MCMC
Introduction

One of the primary goals of modern genetics is to understand the genetic basis of complex traits. What are the genes and alleles that contribute to susceptibility to a particular disease, and how do they interact with each other and with environmental and stochastic factors to produce phenotypes?

The traditional gene mapping approach of positional cloning starts by using linkage analysis in families to identify chromosomal regions that contain genes of interest. These chromosomal regions are typically several centiMorgans in size, and may contain hundreds of genes. Next, linkage analysis is normally followed by linkage disequilibrium and association analysis to help narrow the search down to the functional gene and active variants (e.g. Kerem et al., 1989; Hastbacka et al., 1992).

The positional cloning approach has been very successful at identifying Mendelian genes, but mapping genes for complex traits has turned out to be extremely challenging (Risch, 2000). Despite these difficulties, there has been a mounting number of recent successes in which positional cloning has led to the identification of at-risk haplotypes or occasionally causal mutations, in humans and model organisms (e.g. Horikawa et al., 2000; Laere et al., 2003; Gretarsdottir et al., 2003; Korstanje and Paigen, 2002).

In view of the challenges of detecting genes of small effect using linkage methods, Risch and Merikangas (1996) argued that, under certain assumptions, association mapping is far more powerful than family-based methods. They proposed that in order to unravel the basis of complex traits, the field needed to develop the technical tools for genome-wide association studies (including a genome-wide set of SNPs, and affordable genotyping technology). Those tools are now becoming available, and it will soon be possible to test the efficacy of genome-wide association studies. Moreover, association mapping is already extremely widely used in candidate gene studies (e.g., Lohmueller et al., 2003).

For all these studies, whether or not they start with linkage mapping, association analysis is used to try to detect or localize the active variants at a fine scale. At that point, the data in the LD mapping phase typically consist of genotypes from a subset of the common SNPs in a region. The investigator aims to use these data to detect unobserved variants that impact the trait of interest. For complex traits, it will normally be the case that the active variants have a relatively modest impact on total disease risk. This small signal will be further attenuated if the nearest markers are in only partial LD with the active site (Pritchard and Przeworski, 2001). Moreover, if there are multiple risk alleles in the same gene, these will often arise on different haplotype backgrounds, and may tend to cancel out each other’s signals. (There is a range of views on how serious this problem of allelic heterogeneity is likely to be for complex traits (Terwilliger and Weiss, 1998; Pritchard, 2001; Reich and Lander, 2001; Hugot et al., 2001; Lohmueller et al., 2003).)

For all these reasons, it is important to develop statistical methods that can extract as much information from the data as possible. Certainly, there will be some complex trait loci
that can be detected using very simple analyses. However, by developing more advanced statistical approaches it should be possible to retain power under a wider range of scenarios: e.g., where the signal is rather weak, where the relevant variation is not in strong LD with any single genotyped site (Carlson et al., 2003), or where there is moderate allelic heterogeneity.

Furthermore, for fine mapping, it is vital to use a sensible model to generate the estimated location of disease variants as naive approaches tend to underestimate the uncertainty in the estimates (Morris et al., 2002).

The statistical problem. In this article, we focus on the following problem. Consider a sample of unrelated individuals, each genotyped at a set of markers across a chromosomal region of interest. We assume that the marker spacing is within the typical range of LD, but that it does not exhaustively sample variation. In humans this might correspond to \( \approx 5\) KB spacing on average (Kruglyak, 1999; Zöllner and von Haeseler, 2000; Gabriel et al., 2002). Each individual has been measured for a phenotype of interest, and our ultimate goal is to identify genetic variation that contributes to this phenotype (Figure 1).

With such data, there are two distinct kinds of statistical goals:

1. Testing for association. Do the data provide evidence that there is genetic variation in this region that contributes to the phenotype? (Typically, we would want to see a systematic difference between the genotypes of individuals with high and low phenotype values respectively, or between cases and controls.) The strength of evidence is typically summarized using a p-value.

2. Fine mapping. Assuming that there is variation in this region that impacts the phenotype, then what is the most likely location of the variant(s) and what is the smallest subregion that we are confident contains the variant(s)? This type of information is conveniently summarized as a Bayesian posterior distribution.

The current statistical methods in this field tend to be designed for one goal or the other, but in this article we describe a full multipoint approach for treating both problems in a unified coalescent framework. Our aim is to provide rigorous inference that is more accurate and more robust than existing approaches.

In the first part of this article, we give a brief overview of existing methods for significance testing and fine mapping. Then we describe the general framework of our approach. The middle part outlines our current implementation, developed for case-control data. Finally, we describe results of applications to real and simulated data.

Existing methods.

Significance testing. The simplest approach to significance testing is simply to test each marker separately for association with the phenotype (using a chi-square test of indepen-
Figure 1: Schematic example of the data structure. The lines indicate the chromosomes of three affected individuals (solid) and of three healthy control individuals (dashed). The black circles indicate unobserved variants that increase disease risk. Each column of rectangles indicates the position of a SNP in the dataset. The goal is to use the SNP data to detect the presence of the disease variants and to estimate their location. Notice that for a complex disease we expect to see the “at-risk” alleles at appreciable frequency in controls, and we also expect to find cases without these alleles. As a further complication, there may be multiple disease mutations, each on a different haplotype background.

dence, for example). This approach is most effective when there is a single common disease variant, and less so when there are multiple variants (Slager et al., 2000). When there is a single variant, power is a simple function of $r^2$, the coefficient of LD between the disease variant and the SNP (Pritchard and Przeworski, 2001) and the penetrance of the disease variant. In some recent mapping studies, this simple test has been quite successful (e.g., Van Eerdewegh et al., 2002; Tokuhiro et al., 2003).

The simplest multipoint approach to significance testing is to use two or more adjacent SNPs to define haplotypes, and then test the haplotypes for association (Rioux et al., 2001; Daly et al., 2001; Johnson et al., 2001; Gretarsdottir et al., 2003). It is argued that haplotype-based testing may be more efficient than SNP-based testing at screening for unobserved variants (Johnson et al., 2001; Gabriel et al., 2002). However, there is still uncertainty about how best to implement this type of strategy in a systematic way, and how the resulting power compares to other approaches after multiple-testing corrections.

Various other more complex methods have been proposed for detecting disease association. These include a data mining algorithm (Toivonen et al., 2000), multipoint schemes for identifying identical-by-descent regions in inbred populations (Service et al., 1999; Abney et al., 2002), and for detecting multipoint association in outbred populations (Liang et al., 2001; Tzeng et al., 2003).

Perhaps closest in spirit to the approach taken here is the cladistic approach developed by Alan Templeton and colleagues (Templeton et al. (1987); see also Seltman et al. (2001)).
Their approach is first to construct a set of cladograms based on the marker data by using methods for phylogenetic reconstruction, and then to test whether the cases and controls are non-randomly distributed among the clades. In contrast, the inference scheme presented here is based on a formal population genetic model with recombination. This should enable a more accurate estimation of topology and branch lengths. Our approach also differs from those methods in that we perform a more model-based analysis of the resulting genealogy.

**Fine mapping.** In contrast to the available methods for significance testing, the literature on fine-mapping has a heavier emphasis on model-based methods that consider the genealogical relationships among chromosomes. This probably reflects the view that a formal model is necessary to estimate uncertainty accurately (Morris et al., 2002), and that estimates of location based on simple summary measures of LD do not provide accurate assessments of uncertainty. The challenge is to develop algorithms that are computationally practical, yet extract as much of the signal from the data as possible. The methods should work well for the intermediate penetrance values expected for complex traits, and should be able to deal with allelic heterogeneity.

Though one might ideally wish to perform inference using the ancestral recombination graph (Nordborg, 2001), this turns out to be extremely challenging computationally (e.g., Fearnhead and Donnelly, 2001; Larribe et al., 2002). Instead, most of the existing methods make progress by simplifying the full model in various ways to make the problem more computationally tractable (as we do here).

The first full multipoint, model-based method was developed by McPeek and Strahs (1999). Some elements of their model have been retained in most subsequent models, including ours. Most importantly, they simplified the underlying model by focusing attention only on the ancestry of the chromosomes at each of a series of trial positions for the disease mutation. They then calculated the likelihood of the data at each of those positions, and used the likelihoods to obtain a point estimate and confidence interval for the location of the disease variant. Under that model, non-ancestral sequence could recombine into the data set. The likelihood of non-ancestral sequence was computed using the control allele frequencies, and assuming a first-degree Markov model for the LD between adjacent sites. The McPeek and Strahs model assumed a star-shaped genealogy for the case chromosomes, and applied a correction factor to account for the pairwise correlation of chromosomes due to shared ancestry.

Subsequent variations on this theme have included other methods based on star-shaped genealogies (Morris et al., 2000; Liu et al., 2001), and methods involving bifurcating genealogies of case chromosomes including those of Rannala and Reeve (2001), Morris et al. (2002) and Lam et al. (2000). Two other methods have also used genealogical approaches, but seem to be practical only for very small data sets or numbers of markers (Graham and Thompson, 1998; Larribe et al., 2002). Morris et al. (2002) provide a helpful review of many of these
More recently, Molitor et al. (2003) presented a less model-based multipoint approach to fine mapping. They used ideas from spatial statistics, grouping haplotypes from cases and controls into distinct clusters and assessing evidence for the location of the disease mutation from the distribution of cases across the clusters. Their approach may be more computationally feasible for large data sets than are fully model-based genealogical methods, but it is unclear if some precision is lost by not using a coalescent model.

The procedure described in the present differs from existing methods in several important aspects. Our approach estimates the joint genealogy of all individuals, not just of cases. This should allow us to model the ancestry of the sample more accurately, and to include allelic heterogeneity in a more realistic way. We also analyze the evidence for the presence of a disease mutation after inferring the ancestry of a locus. This enables us to apply realistic models of penetrance and to analyze quantitative traits. Furthermore, in our Markov chain algorithm we do not record the full ancestral sequences at every node, which should enable better mixing and allow analysis of larger datasets.

Models and Methods

We consider the situation where the data consist of a sample of individuals who have been genotyped at a set of markers spaced across a region of interest (Figure 1). Each individual has been assessed for some phenotype, which can be either binary (e.g., affected with a disease, or unaffected) or quantitative. Our framework can also accommodate TDT data (Spielman et al., 1993), where the untransmitted genotypes are treated as controls.

We are most interested in the setting where the genotyped markers represent only a small fraction of the variation in the region, and our goal is to use LD and association to detect unobserved susceptibility variants. We allow for the possibility of allelic heterogeneity (there might be multiple independent mutation events that produce susceptibility alleles), but we assume that all these mutations occur close enough together (e.g., within a few kb) that we can treat them as having a single location within the region.

The genealogical approach. The underlying model for our approach is derived from the coalescent (reviewed by Hudson, 1990; Nordborg, 2001). The coalescent refers to the conceptual idea of tracing the ancestry of a sample of chromosomes back in time. Even chromosomes from “unrelated” individuals in a population share a common ancestor at some time in the past. Moving backward in time, eventually all the lineages that are ancestral to a modern day sample “coalesce” to a single common ancestor. The time-scales for this process are typically rather long—for example, the most recent common ancestor of human \( \beta \)-globin sequences is estimated to have been about 800,000 years ago (Harding et al., 1997).

When there is recombination, the ancestral relationships among chromosomes are more
Figure 2: Hypothetical example of a coalescent genealogy for a sample of 28 chromosomes, at the locus of a disease susceptibility gene. Each tip at the bottom of the tree represents a sampled chromosome; the lines indicate the ancestral relationships among the chromosomes. The two black circles on the tree represent two independent mutation events producing susceptibility variants. These are inherited by the chromosomes marked with gray filled circles. Individuals carrying those chromosomes will be at increased risk of disease. This means that there will be a tendency for chromosomes from affected individuals to cluster together on the tree, in two mutation-carrying clades. The degree of clustering depends in part on the penetrance of the mutation.

complicated. At any single position along the sequence, there is still a single tree, but the trees at nearby positions may differ. It is possible to represent the full ancestral relationships among chromosomes using a concept known as the “ancestral recombination graph” (ARG) (Nordborg, 2001; Nordborg and Tavare, 2002), although it is difficult to visualize the ARG except for small samples or short chromosomal regions (Figure 3).

Considering the coalescent process provides useful insight into the nature of the information about association that is contained in the data. Figure 2 shows a hypothetical example of the coalescent ancestry of a sample of chromosomes at the position of a disease susceptibility locus. In this example, there are two disease susceptibility mutations present in the sample. By definition, these will be carried at a higher rate in affected individuals than in controls. This implies that chromosomes from affected individuals will tend to be non-randomly clustered on the tree. Each independent disease mutation gives rise to one cluster of “affected” chromosomes.

Traditional methods of association mapping work by testing for association between the phenotype status and alleles at linked marker loci (or with haplotypes). In effect, association at a marker indicates that in the neighborhood of this marker, chromosomes from affected individuals are more closely related to one another than random. Fundamentally, the marker data are informative because they provide indirect information about the ancestry of un-
observed disease variants. Detecting association at non-causative SNPs implies that case chromosomes are non-randomly clustered on the tree.

In fact, unless we have the actual disease variants in our marker set, the best information that we could possibly get about association is to know the full coalescent genealogy of our sample at that position. If we knew this, the marker genotypes would provide no extra information; all the information about association is contained in the genealogy.

Hence, our approach is to use the marker information to learn as much as we can about the coalescent genealogy of the sample at different points along the chromosome. Our statistical inference for association- or fine-mapping will be based on this. In what follows, we outline our approach of using marker data to estimate the unknown coalescent ancestry of a sample, and describe how this information can be used to perform inference. Unlike previous mapping methods (e.g., Morris et al., 2002), we aim to reconstruct the genealogy of the entire sample, and not just the genealogy of cases. This extension allows us to extract substantially more information from the data, and enables significance testing.

Performing inference. We start by developing some notation. Consider a sample of $n$ haplotypes from $n/2$ unrelated individuals. The phenotype of individual $i$ is $\phi_i$, and $\Phi$ represents the vector of phenotype data for the full sample of $n/2$ individuals. The phenotypes might be qualitative (e.g., affected/unaffected), or quantitative measurements.

Each individual is genotyped at a series of marker loci from one or more genomic regions (or in the future possibly from genome-wide scans). Let $G$ denote the multidimensional vector of haplotype data—i.e., the genotypes for $n$ haplotypes at $L$ loci (possibly with missing data). Let $X$ be the set of possible locations of the QTL or disease susceptibility gene and let $x \in X$ represent its (unknown) position. Our approach will be to scan sequentially across the regions containing genotype data, considering many possible positions for $x$. A natural measure of support for the presence of a disease mutation at position $x$ is given by the likelihood ratio (LR):

$$LR = \frac{L_A(\Phi; x, \hat{P}_{alt}, G)}{L_0(\Phi; \hat{P}_0, G)},$$

where $L_A$ and $L_0$ represent likelihoods under the alternative model (disease mutation at $x$), and null hypothesis (no disease mutation in the region), respectively. $\hat{P}_{alt}$ and $\hat{P}_0$ are the vectors of penetrance parameters under the alternative and null hypotheses, respectively, that maximize the likelihoods. Large values of the likelihood ratio indicate that the null hypothesis should be rejected. Specific models to calculate these likelihoods are described below (see equations 6 and 7).

We also want to estimate the location of disease mutations. For this purpose it is convenient to adopt a Bayesian framework, as this makes it more straightforward to account for the various sources of uncertainty in a coherent way (Morris et al., 2000; Liu et al., 2001;
Morris et al., 2002). The posterior probability that a disease mutation is at $x$ is then

$$
Pr(x \mid \Phi, G) = \frac{Pr(\Phi, G \mid x) Pr(x)}{\int_x Pr(\Phi, G \mid y) Pr(y) dy} \propto Pr(\Phi, G \mid x) Pr(x)
$$

(2)

where $Pr(x)$ gives the prior probability that the disease locus is at $x$. $Pr(x)$ will normally be set uniform across the genotyped regions, but this prior can easily be modified to take advantage of prior genomic information if desired (see discussion in Rannala and Reeve, 2001; Morris et al., 2002).

In order to evaluate expressions (1) and (2), we need to compute $Pr(\Phi, G \mid x)$. In order to do so, we introduce the notation $T_x$, to represent the (unknown) coalescent genealogy of the sample at $x$. $T_x$ records both the topology of the ancestral relationships among the sampled chromosomes and the times at each internal node. Then

$$
Pr(\Phi, G \mid x) = \int Pr(\Phi, G \mid x, T_x) Pr(T_x \mid x) dT_x
$$

(3)

where the integral is evaluated over all possible trees. We now make the following approximations: (i) $Pr(T_x \mid x) \approx Pr(T_x)$ and (ii) $Pr(G \mid \Phi, x, T_x) \approx Pr(G \mid T_x)$. The first approximation implies that in the absence of the phenotype data, the tree topology itself contains no information about the location of the disease mutation. Thus, we ignore the possible impact of selection and over-ascertainment of affected individuals in changing the distribution of branch times at the disease locus. Our expectation is that the data will be strong enough to overcome minor mis-specification of the model in this respect (this was the experience of Morris et al., 2002, in a similar situation). The second approximation is a good assumption if the active disease mutation is not actually in our marker set and if mutations at different positions occur independently. We can then write

$$
Pr(\Phi, G \mid x) \approx \int Pr(\Phi \mid x, T_x) Pr(G \mid T_x) Pr(T_x) dT_x
$$

and since $Pr(G \mid T_x)Pr(T_x) = Pr(T_x \mid G)Pr(G)$ we obtain

$$
Pr(\Phi, G \mid x) \approx \int Pr(\Phi \mid x, T_x) Pr(T_x \mid G) Pr(G) dT_x
$$

$$
\propto \int Pr(\Phi \mid x, T_x) Pr(T_x \mid G) dT_x.
$$

(4)

Expression (4) consists of two parts. $Pr(\Phi \mid x, T_x)$ is the probability of the phenotype data given the tree at $x$. To compute this, we specify a disease model, and then integrate over the possible branch locations of disease mutations in the tree (see below for details). $Pr(T_x \mid G)$ refers to the posterior density of trees given the marker data and a population genetic model.
to be specified; the next subsection outlines our approach to drawing Monte Carlo samples from this density.

In summary, our approach is to scan sequentially across the region(s) of interest, considering a dense set of possible positions of the disease location \( x \). At each position \( x \), we sample \( M \) trees (denoted \( T_x^{(m)} \)) from the posterior distribution of trees. For Bayesian inference of location, we apply Equation 2 to estimate the posterior density \( \Pr(x \mid \Phi, G) \) at \( x \) by computing

\[
\Pr(x \mid \Phi, G) \approx \frac{\frac{1}{M} \sum_{m=1}^{M} \Pr(\Phi \mid x, T_x^{(m)}) \Pr(x)}{\sum_{i=1}^{Y} \frac{1}{M} \sum_{m=1}^{M} \Pr(\Phi \mid y_i, T_{yi}^{(m)}) \Pr(y_i)},
\]

where \( \{y_1, \ldots, y_Y\} \) denote a series of \( Y \) trial values of \( x \) spaced across the region of interest. We will occasionally refer to the numerator of Equation 5, divided by \( \Pr(x) \), as the “average posterior likelihood” at \( x \). For significance testing at \( x \), we maximize

\[
L_A(\Phi; x, \hat{P}_{alt}, G) \approx \frac{1}{M} \sum_{m=1}^{M} \Pr(\Phi \mid x, T_x^{(m)}, \hat{P}_{alt})
\]

and

\[
L_0(\Phi; \hat{P}_0, G) = \Pr(\Phi \mid \hat{P}_0)
\]

with respect to \( \hat{P}_{alt} \) and \( \hat{P}_0 \). See below for details about how these probabilities are computed.

**Sampling from the genealogy, \( T_x \).** To perform these calculations, it is necessary to sample from the posterior density \( T_x \mid G \) (loosely speaking, we wish to draw from the set of coalescent genealogies that are consistent with the genotype data). We adopt a fairly standard population genetic model, namely the neutral coalescent with recombination (i.e., the ancestral recombination graph (ARG); Nordborg, 2001). Our current implementation assumes constant population size.

A number of recent studies have aimed to perform full likelihood or Bayesian inference under the ARG (Griffiths and Marjoram (1996); Fearnhead and Donnelly (2001); Nielsen (2000); Kuhner et al. (2000); Larribe et al. (2002); reviewed by Stephens (2001)). The experience of these earlier studies indicates that this is a technically challenging problem, and that existing methods tend to perform well only for quite small data sets (e.g., Wall, 2000; Fearnhead and Donnelly, 2001). Therefore, we have decided to perform inference under a simpler, local approximation to the ARG, reasoning that this might allow accurate inference for much larger datasets. Our implementation applies Markov chain Monte Carlo (MCMC) techniques (see Appendix 1).

In our approximation, we aim to reconstruct the coalescent tree only at a single “focal point” \( x \), though we use the full genotype data from the entire region, as all of this is potentially informative about the tree at that focal point. Consider two chromosomes that have a very recent common ancestor (at the focal point). These chromosomes will normally both inherit a large region of chromosome around the focal point, uninterrupted by recombination,
Figure 3: Hypothetical example of the ancestral recombination graph (ARG) for a sample of 6 chromosomes, labeled A-F (left plot), along with our representation (middle and right plots). Left plot: The ARG contains the full information about the ancestral relationships among a sample of chromosomes. Moving up the tree from the bottom (backwards in time), points where branches join indicate coalescent events, while splitting branches represent recombination events. At each split, a number indicates the position of the recombination event (for concreteness, we assume nine inter-marker intervals, labeled 1–9). By convention, the genetic material to the left of the breakpoint is assigned to the left branch at a split. See Nordborg (2001) for a more extensive description of the ARG. Middle and right plots: At each point along the sequence, it is possible to extract a single genealogy from the ARG. The plots show these genealogies at two “focal points”, located in intervals 4, and 7, respectively. The numbers in parentheses indicate the total region of sequence that is inherited without recombination, along with the focal point, by at least one descendant chromosome. (1,9) indicates inheritance of the entire region. For clarity, not all intervals with complete inheritance (1,9) are shown.
from that one common ancestor. Then consider a more distant ancestor that the two chromosomes share with a third chromosome in the sample. It is likely that the region around the focal point shared by the three chromosomes is smaller. In our representation of the genealogy, we store the topology at the focal point, along with the extent of sequence at each node that is ancestral to at least one of the sampled chromosomes without recombination (Figure 3).

An example of this is provided in Figure 4. Each tip of the tree records the full sequence (across the entire region) of one observed haplotype. Then, moving up the tree, as the result of a recombination event a part of the sequence may split off and evolve on a different branch of the ARG. When this happens, the amount of sequence that is co-evolving with the focal point is reduced. The length of the sequence fragment that co-evolves with the focal point can increase during a coalescent event, as the sequence in the resulting node is the union of the two coalescing sequences. In other words, the amount of sequence surrounding the focal point shrinks when a recombination event occurs and may increase at a coalescent event. A marker is retained up to a particular node as long as there is at least one lineage leading to this node in which that SNP is not separated from the focal point by recombination. We do not model coalescent events in the ARG where only one of the two lines carries the focal point. Therefore, the sequence at internal nodes will always consist of one contiguous fragment of sequence.

Our MCMC implementation stores the tree topology, node times, and the ancestral sequence at each node. We assume a finite sites mutation model for the markers that are retained on each branch. (This rather simplistic model is far more computationally convenient than more realistic alternatives.) At some points, sequence is introduced into the genealogy through recombination events. We approximate the probability for the introduced sequence by assuming a simple Markov model based on the allele frequencies in the sample (similar approximations have been used previously by McPeek and Strahs, 1999; Morris et al., 2000; Liu et al., 2001; Morris et al., 2002). The population recombination rate $\rho$, and the mutation rate $\theta$, are generally unknown in advance, and are estimated from the data within the MCMC scheme, assuming uniform rates along the sequence. A more precise specification of the model and algorithms is provided in Appendix 1.

Overall, our model is similar to those of earlier approaches such as the haplotype sharing model of McPeek and Strahs (1999) and the coalescent model of Morris et al. (2002). However, we focus on chromosomal sharing backwards in time, rather than decay of sharing from an ancestral haplotype. In part, this reflects our shift away from modeling affected chromosomes only, to modeling the tree for all chromosomes. The representation used by those earlier studies means that they potentially have to sum over possible ancestral genotypes at sites far away from the focal point $x$, that are not ancestral to any of the sampled chromosomes and about which there is therefore no information. Storing all this extra information is likely to be detrimental in an MCMC scheme, as it presumably impedes rearrangements of
Figure 4: Example of an ancestral genealogy as modeled by our treebuilding algorithm. The figure shows the ancestry of a single focal point (designated F) as inferred from three biallelic markers (alleles are shown as 0 and 1). Branches with recombination events on them are depicted as red lines, showing at the tip of the arrow the part of the sequence that evolves on a different genealogy. As can be seen at the coalescent event at time $t_4$, if no recombination occurs on either branch, the entire sequence is transmitted along a branch and coalesces, generating a full-length sequence. If on the other hand a recombination event occurs, the amount of sequence that reaches the coalescent event is reduced (indicated by the dashes). If this reduction occurs on only one of the two branches, the sequence can be restored from the information on the other branch (as at time $t_3$). But if recombination events occur on both branches, the length of the sequence is reduced ($t_2$).
the topology. Thus, we believe that our representation can potentially improve both MCMC mixing and the computational burden involved in each update. Indeed, if one wished to perform inference across an infinitely long chromosomal region, the total amount of sequence stored at the ancestral nodes in our representation would be finite, while that in the earlier methods would not.

A more fundamental difference is that, unlike most of the previous model-based approaches to this problem, our genealogical reconstruction is independent of the phenotype data. There are trade-offs in choosing to frame the problem in this way, as follows. When the alternative model is true, the phenotype data contain some information about the topology which could help to guide the search through tree-space. In contrast, our procedure weights the trees after sampling them from \( \Pr(T_x \mid G) \) according to how consistent they are with the phenotype data (Eqns. 5-6), ignoring additional information from the phenotype data. However, tackling the problem in this way makes it far easier to assess significance, because we know that under the null the phenotypes are randomly distributed among tips of the tree. It also means that we can calculate posterior densities for multiple disease models using a single MCMC run.

**Modeling the phenotypes.** In order to compute expressions (5) and (6), we use the following model to evaluate \( \Pr(\Phi \mid x, T_x) \). At the unobserved disease locus, let \( A \) denote the genotype at the root of the tree \( T_x \). We assume that genotype \( A \) mutates to genotype \( a \) at rate \( \nu/2 \) per unit time, independently on each branch. We further assume that alleles in state \( a \) do not undergo further mutation.

Next, we need to define a model for the genotype-phenotype relationship for each of the three diploid genotypes at the susceptibility locus: namely, \( \Pr(\phi \mid AA) \), \( \Pr(\phi \mid Aa) \), and \( \Pr(\phi \mid aa) \), where \( \phi \) refers to a particular phenotype value (e.g., affected/unaffected, or a quantitative measure). For a binary trait, these three probabilities simply denote the genotypic penetrances: e.g., \( \Pr(\text{Affected} \mid AA) \). In practice, the situation is often complicated by the fact that the sampled individuals may not be randomly ascertained. In that case, the estimated “penetrances” really correspond to \( \Pr(\phi \mid AA, S) \), \( \Pr(\phi \mid Aa, S) \), and \( \Pr(\phi \mid aa, S) \), where \( S \) refers to some sampling scheme (e.g., choosing equal numbers of cases and controls).

In the algorithm presented here, we assume that the affection status of the two chromosomes in an individual can be treated independently from each other and from the frequency of the disease mutation: i.e., \( P_A(\phi) \) is the probability that a chromosome with genotype \( A \) comes from an individual with phenotype \( \phi \), and analogously for \( P_a(\phi) \). In the binary situation, this model has two independent parameters: \( P_A(1) = 1 - P_A(0) \) and \( P_a(1) = 1 - P_a(0) \). In this case the ratio \( \frac{P_A(\phi)}{P_a(\phi)} \) corresponds directly to the relative risk of allele \( A \), conditional on the sampling scheme. As another example, for a normally distributed trait, \( P_A(\phi) \) and \( P_a(\phi) \) are the densities of two normal distributions at \( \phi \), and would be characterized by mean and variance parameters. Note that most values of \( P_A(\phi) \) and \( P_a(\phi) \) do not correspond to a single
genetic model exists as the mapping from \((P_A(\phi), P_a(\phi))\) to \((\Pr(\phi|AA), \Pr(\phi|Aa), \Pr(\phi|aa))\) is dependent on the frequency of the disease mutation. Nevertheless, this factorization of the penetrance parameters is computationally convenient and allows for an efficient analysis of \(T_x\).

Of course, it is not known in advance which chromosomes are \(A\) and which are \(a\), so we compute the likelihood of the phenotype data by summing over the possible arrangements of mutations at the disease locus. Under the alternative hypothesis, most arrangements of mutations will be relatively unsupported by the data, while branches leading to clusters of affected chromosomes will have high support for containing mutations. Let \(M\) record which branches of the tree contain disease mutations and \(\gamma \subset \{1, \ldots, n\}\) be the set of chromosomes that carry a disease mutation according to \(M\) (i.e. the descendants of \(M\)) and let \(\beta\) be the set of chromosomes that do not carry a mutation, i.e., \(\beta = \{1, \ldots, n\} \setminus \gamma\). Then we calculate

\[
\Pr(\Phi | x, T_x, \nu) = \sum_M \left( \prod_{i \in \gamma} P_A(\phi_i) \prod_{i \in \beta} P_a(\phi_i) \Pr(\gamma | x, T_x, \nu) \right)
\]

(8)

For a case-control dataset this can be written as

\[
\Pr(\Phi | x, T_x, \nu) = \sum_M P_A^{n^A}(1 - P_A)^{n^A} P_a^{n^a}(1 - P_a)^{n^a} \Pr(M | x, T_x, \nu)
\]

where \(n^A_i\) and \(n^a_i\) count the number of \(i\)-type chromosomes (where \(i \in \{A, a\}\)) from affected and healthy individuals, respectively. Equation (8) can be evaluated efficiently using a peeling algorithm (Felsenstein, 1981). The details of this algorithm are provided in Appendix 2. Calculations for general diploid penetrance models are much more computationally intensive, and we will present those elsewhere.

For our Bayesian analysis, we take the prior for the parameters governing \(P_A(\phi)\) and \(P_a(\phi)\) to be uniform and independent on a bounded set \(\Delta\), and average the likelihoods over this prior. By allowing any possible order for the penetrances under the alternative model, we allow for the possibility that the ancestral allele may actually be the high-risk allele, as observed at some human disease loci, including ApoE (Fullerton et al., 2000).

For significance testing, we test the null hypothesis that \(P_A(\phi) = P_a(\phi)\) compared to the alternative model where the parameters governing \(P_A(\phi)\) and \(P_a(\phi)\) can take on any values independently. Standard theory suggests that twice the log likelihood ratio of the alternative model, compared to the null, should be asymptotically distributed as \(\chi^2\) random values with \(d\) degrees of freedom, where \(d\) is the number of extra parameters in the alternative model compared with the null. Thus, for case-control studies our formulation suggests that twice the log likelihood ratio should have a \(\chi^2\) distribution. In fact, simulations that we have done (results not shown) indicate that this assumption is somewhat conservative.

Lastly, it remains to determine the mutation rate, \(\nu\), at the unobserved disease locus. It seems unlikely that there will be much information in the data about \(\nu\), hence we prefer to set it to a plausible value, \(a\) priori. For a similar model, Pritchard (2001) argued that the
most biologically plausible values for this parameter are in the range of about 0.1 to 1.0, corresponding to low, and moderate levels of allelic heterogeneity, respectively.

**Multiple testing.** Typically, association mapping studies consist of large numbers of statistical tests. To account for this, it is common practice to report a p-value that measures the significance of the largest departure from the null hypothesis anywhere in the dataset. The simplest approach is to apply a Bonferroni correction (i.e., multiplying the p-value by the number of tests), but this tends to be unnecessarily conservative because the association tests at neighboring positions are correlated.

A more appealing solution is to use randomization techniques to obtain an empirical overall p-value (c.f. McIntyre et al., 2000). The basic idea is to hold all the genotype data constant, and randomly permute the phenotype labels. For each permuted set, the tests of association are repeated, and the smallest p-value for that set is recorded. Then the experiment-wide significance of an observed p-value \( p_i \) is estimated by the fraction of random datasets whose smallest value is \( \leq p_i \).

The latter procedure is only practical if the test of association is computationally fast. For the method proposed in this paper, the inference of ancestries is independent of phenotypes. Therefore, the trees need to be generated only once in this scheme and the sampled trees are stored in computer memory. Then, the likelihood calculations can be performed on these trees using both the real, and randomized phenotype data to obtain the appropriate empirical distribution. For a whole genome scan, a permutation test with the proposed peeling strategy is rather daunting. Performing the peeling analysis for 1000 permutations on one tree of 100 cases and 100 controls takes about 6 minutes on a modern desktop machine. Thus, a whole genome permutation test with one focal point every 50 kb, 100 trees per focal point and a penetrance grid of \( 19 \times 19 \) values would take about 750,000 processor hours.

A rather different solution for genome-wide scans of association may be to apply the false discovery rate criterion, as this tends to be robust to local correlation when there are enough independent data (Benjamini and Hochberg, 1995; Sabatti et al., 2003).

**Unknown haplotype phase.** Our current implementation assumes that the individual genotype data can be resolved into haplotypes. However, in many current studies, haplotypes are not experimentally determined, and must instead be estimated by statistical methods. In principle, it would be natural to update the unknown haplotype phase within our MCMC coalescent framework described below (Lu et al., 2003; Morris et al., 2003). By doing so, we would properly account for the impact of haplotype uncertainty on the analysis. In fact, Morris et al. (2004) concluded that doing so increased the accuracy of their fine-mapping algorithm (compared to the answers obtained after estimating haplotypes via a rather simple EM procedure). However, it is already a difficult problem to sample adequately from the posterior distribution of trees given known haplotypes and it is unclear to us that the added
burden of estimating haplotypes within the MCMC scheme represents a sensible tradeoff. Therefore, we currently use point estimates of the haplotypes obtained from PHASE 2.0 (Stephens et al., 2001; Stephens and Donnelly, 2003). We also currently use PHASE to impute missing genotypes.

**False positives due to population structure.** It has long been known that case-control studies of association are susceptible to high Type 1 error rates when the samples are drawn from structured or admixed populations (Lander and Schork, 1994). Therefore, we advise using unlinked markers to detect problems of population structure (Pritchard and Rosenberg, 1999), prior to using the association mapping methods presented here.

When population structure is problematic, there are two types of methods that aim to correct for it: Genomic Control (Devlin and Roeder, 1999) and Structured Association (Pritchard et al., 2000; Satten et al., 2001). It seems likely that some form of Genomic Control correction might apply to our new tests, but it is not clear to us how to obtain this correction theoretically. It should be possible to obtain robust p-values using the Structured Association approach roughly as follows. First, one would apply a clustering method to the unlinked markers to estimate the ancestry of the sampled individuals (Pritchard et al., 2000; Satten et al., 2001), and the phenotype frequencies across subpopulations. Then, the phenotype labels could be randomly permuted across individuals while preserving the overall phenotype frequencies within subpopulations. As before, the test statistic of interest would be computed for each permutation.

**SNP ascertainment and heterogeneous recombination rates.** In the MCMC algorithm described above, and more fully in the Appendix, we assume—for convenience—that mutation at the markers can be described using a standard finite sites mutation model with mutation parameter $\theta$. However in practice, we aim to apply our method to SNPs: markers for which the mutation rate per site is likely to be very low, but which have been specifically ascertained as polymorphic. Hence, our estimate of $\theta$ should not be viewed as an estimate of the neutral mutation rate; it is more likely to be roughly the inverse of the expected tree length (if there has usually been one mutation per SNP in the history of the sample). Moreover, the facts that SNPs are often ascertained to have intermediate frequency, and that we over-estimate $\theta$ may lead to some distortion in the estimated branch lengths. However, we anticipate that most of the information about the presence of disease variation will come from the degree to which case and control chromosomes cluster on the tree, so bias in the branch length estimates may not have a serious impact on inferences about the location of disease variation. The next section provides results supporting this view.

Another factor not considered in our current implementation is the possibility of variable recombination rate (e.g., Jeffreys et al., 2001). Since recombination rates appear to vary considerably over quite fine scales, this is probably an important biological feature to
include in analysis. One route forward would be for us to estimate separate recombination parameters in each inter-marker interval, within the MCMC scheme (perhaps correlated across neighboring intervals). It is unclear how much this would add to the computational burden of convergence and mixing. In the short term, it would be possible to use a separate computational method to estimate these rates prior to analysis with LATAG (e.g., using Li and Stephens, 2003), and to modify the input file to reflect the estimated genetic distances.

Software. The algorithms presented here have been implemented in a program called “LATAG” (Local Approximation to the Ancestral recombination Graph). The program is available on request from SZ.

Testing and Applications: Simulated data

Simulation methods. To provide a systematic assessment of our algorithm we simulated 50 datasets, each representing a fine mapping study or a test for association within a candidate region. Each dataset consists of 30 diploid cases and 30 diploid controls that have been genotyped for a set of markers across a region of 1 cM. Our model corresponds to a scenario of a complex disease locus with relatively large penetrance differences (since the sample sizes are small), and with moderate allelic heterogeneity at the disease locus.

The datasets were generated as follows. We simulated the ancestral recombination graph (ARG), assuming a constant population size of 10,000 diploid individuals, and a uniform recombination rate. On the branches of this ARG, mutations occurred as a Poisson process according to the infinite sites model. The mutation rate was set so that in typical realizations there would be 45-65 markers with minor allele frequency > 0.1 across the 1 cM region. The position of the disease locus $x_s$ was drawn from a uniform distribution across the region. Mutation events at the disease locus were simulated on the tree at that location at rate 1 per unit branch length (in coalescent time), with no back mutations (c.f. Pritchard, 2001). This process determines whether each chromosome does, or does not, carry a disease mutation. We required that the total frequency of mutation-bearing chromosomes be in the range 0.1-0.2, and if it was not, then we simulated a new set of disease mutations at the same location. This procedure generated a total of 10-25 disease mutations across the entire population, though many of the mutations were redundant or at low frequency.

In order to assign phenotypes, we the following penetrances: a homozygote wild-type showed the disease phenotype with probability $P_{hw} = 0.05$, a heterozygous genotype with probability $P_{he} = 0.1$, and a homozygous mutant with probability $P_{hm} = 0.8$. According to these penetrances, we then created 30 case and 30 control individuals by sampling without replacement from the simulated population of 20,000 chromosomes, as follows. Let $n$ be the remaining number of wild-type chromosomes in the population and $m$ the remaining number of mutant chromosomes in the population. Then the next case individual was homozygous for
the mutation with probability 
\( P_{hm} m \cdot (m-1) \cdot (P_{hm} m \cdot (m-1) + 2 \cdot P_{he} m \cdot n + P_{hw} n \cdot (n-1))^{-1}, \)

heterozygous with probability 
\( (2 \cdot P_{he} m \cdot n) \cdot (P_{hm} m \cdot (m-1) + 2 \cdot P_{he} m \cdot n + P_{hw} n \cdot (n-1))^{-1}, \)

and otherwise homozygous for the mutant allele. The diplotypes for each case were then
created by sampling the corresponding number of mutant or wildtype chromosomes. Control
individuals were generated analogously. Across the 50 replicates, we found that 10-33 of the
60 case chromosomes and 0-9 of the control chromosomes carried a disease mutation.

As might be expected for the simulation of a complex disease, not all of the simulated
datasets carried much information about the presence of genetic variation influencing the
phenotype. For instance, in 22 of the generated datasets, the highest single point association
signal among the generated markers, calculated as Pearson’s \( \chi^2 \), is below 6.5.

We analyzed each simulated dataset by considering 50 focal points \( x_1, \ldots, x_{50} \), spaced
equally across the 1 cM region. For each point \( x_i \) we used LATAG to draw 50 trees from
the distribution \( Pr(T_{x_i} | G, x_i) \). To ensure convergence of the MCMC, we used a burn-in
period of \( 2.5 \cdot 10^6 \) iterations for \( x_1 \). As the tree at location \( x_i \) is a good starting guess for
trees of the adjacent tree at \( x_{i+1} \) we used a burn-in of \( 0.5 \cdot 10^6 \) iterations for \( x_2, \ldots, x_k \).
We sampled each set of trees \( \{T_{x_1}^{(1)}, \ldots, T_{x_50}^{(50)}\} \) using a thinning interval of 10,000 steps and
estimated \( Pr(\Phi, G | x_i) \) according to (5) and (18) without assuming any prior information
about the location of disease mutations. We found that the mean was somewhat unstable
due to occasional large outliers, and therefore substituted the median for the average in
(5). To evaluate (18) we summed over a grid of penetrances \( \Delta = \{0.05, 0.1, \ldots, 0.95\} \times \\
\{0.05, 0.1, \ldots, 0.95\} \), setting the disease mutation rate \( \nu \) to 1.0. We calculated the posterior
probability at each locus \( x_i \) by evaluating

\[
Pr(x_i | \Phi, G) = \frac{Pr(\Phi, G | x_i)}{\sum_{j=1}^{50} Pr(\Phi, G | x_j)}.
\]

In addition, we recorded the point estimate for the location of the disease mutation as
the \( x_i \) with the highest posterior probability. The running time for each dataset was about
5 hours on a 2.4 Ghz processor with 512 K memory.

For comparison, we also analyzed each dataset with DHSMP map 2.0 using the standard
settings suggested in the program package. This program generated point estimates for the
locus of disease mutation and two 95% confidence intervals: the first assuming a star-like
phylogeny among cases, and the second using a correction to account for the additional
correlation among cases that results from relatedness.

Significance tests were performed by two methods. First we calculated

\[
L_m = \max\{L_A(\Phi, x_i, P_{alt}, G) : i \in \{1, \ldots, 50\}, \ P_{alt} \in \Delta\}
\]

(9)

and calculated the likelihood ratio according to (1):

\[
LR = \frac{L_m}{L_0}.
\]
with \( L_0 = 0.5^{120} \). We assigned pointwise significance to this ratio by assuming that \( 2 \cdot \ln(LR) \) is \( \chi^2 \)-distributed with one degree of freedom. (Other simulations that we have done indicate that this assumption is somewhat conservative; results not shown.) To estimate global significance, we permuted case and control status among the 60 individuals 1000 times, recalculated \( L_m \) for each permutation (using the original trees obtained from the data) and counted the number of permutations that showed a higher \( L_m \) than the original dataset anywhere in the region. The permutation procedure corrects for multiple testing across the region, and does not rely on the predicted distribution of the likelihood ratio.

For comparison we assessed the performance of single-point association analysis by calculating the association of each observed marker in a \( 2 \times 2 \)-contingency table with a \( \chi^2 \) statistic and recorded the \( \chi^2 \) of the marker with the highest value. We assigned significance to this test statistic in two ways: first based on the \( \chi^2 \)-distribution with one degree of freedom, and second, by performing 1000 permutations of phenotypes among the 60 individuals and counting the number of permutations in which the highest observed \( \chi^2 \) was higher than observed in the sample.

In order to assess convergence, we then repeated the analysis of each dataset an additional four times and compared the estimated posterior distributions to assess the convergence of the MCMC and the variability in estimation. We calculated the overlap of two credible intervals \( C_1 \) and \( C_2 \) obtained from multiple MCMC analyses of the same dataset as

\[
\left( \frac{|C_1 \cap C_2|}{|C_1|} + \frac{|C_1 \cap C_2|}{|C_2|} \right) \div 2
\]

where \(|I|\) is the length of interval \( I \).

**Results**

**Assessing convergence.** An important issue for MCMC applications is to check the convergence of the Markov chain, since poor convergence or poor mixing can lead to unreliable results. While numerous methods exist to diagnose MCMC performance, (Gammerman, 1997), the most direct approach is to compare the results from multiple MCMC runs. If the Markov chain performs well, and the samples drawn from the posterior are sufficiently large, then different runs will produce similar results. (Conversely, good performance by this criterion does not absolutely guarantee that the Markov chain is working well, but it is certainly encouraging.)

To assess the convergence of the LATAG algorithm, we performed 5 runs for each of the 50 simulated datasets. For our simulated datasets we found that on average, pairs of 50% and credible intervals overlapped by 75% and pairs of 95% credible intervals overlapped by 96%.

As a second method of evaluating the convergence of the MCMC, we compared the point estimates for the location of disease mutation between the different runs. The average distance between two point estimates on the same dataset is 184 kb. This number includes
datasets where there is very little information about the locus of disease mutation. Figure 5 displays the point estimates for across independent runs, indicating that for most datasets all five runs produce a similar estimate. Further inspection of the results in Figure 5 indicates that in most cases where there is substantial variation across runs, this is because the posterior distribution is relatively flat across the entire region. Some of the datasets contain very little information about the presence or location of disease mutations, and so small random fluctuations in the estimation can shift the peak from one part of the region to another. To further quantify this observation, we computed the correlation between the average pairwise difference of the point estimates with the average posterior likelihood at the point estimate for each dataset. We observed that these were strongly negatively correlated (correlation coefficient = -0.29). The higher the signal that is present in the data (expressed in posterior probability), the smaller the difference between the point estimates.

In summary, when the datasets contained a strong signal, the concordance between individual runs was quite high, indicating good convergence. On the other hand, when the information about location was weak, random variation across runs meant that the point estimates sometimes varied considerably. In such cases, longer runs would be needed to obtain really accurate estimates. For analyzing real data, it is certainly important to use multiple LATAG runs to ensure the robustness of the results.
Point estimates of location. The mode of the posterior distribution is a natural “best guess” for the location of the disease variation. To assess the accuracy of this estimate, we calculated the distance between this point estimate and the real locus of the disease mutation for each simulated dataset. Overall, we observed an average error of 0.19 cM with a standard deviation of 0.23 cM. To evaluate this result, we compared it to the accuracy of two other point estimators. As a naive estimator, we chose the position of the marker that has the highest level of association with the phenotype, measured using the Pearson $\chi^2$ statistic. This choice is based on the observation that, on average, LD declines with distance. As an example of an estimator provided by a multipoint method, we analyzed the prediction generated by DHSMAP.

We found that the average distance between the disease locus and the SNP with the highest $\chi^2$ was 0.25 cM (standard deviation 0.26 cM) and the distance to the DHSMAP point estimate was 0.27 cM (standard deviation 0.25 cM). The cumulative distributions of the error in estimation are displayed in Figure 6. The estimate generated by LATAG is most likely to be close to the real locus of disease mutation. For instance, in 54% of all simulations, the LATAG estimate is within 0.1 cM of the real locus, while the naive estimate is within 0.1 cM in 44% of all cases and DHSMAP is in the same range in 30% of our simulated datasets.
Figure 7: Coverage accuracy of the credible intervals obtained by LATAG. We constructed credible intervals of different sizes, ranging from 10% to 90% (x-axis). The y-axis shows the number of datasets (out of 50) for which the credible interval contained the true location of the disease gene. The blue bars (number observed) are generally slightly higher than the red bars (number expected if the coverage is correct), suggesting that our credible intervals may be slightly conservative.

Coverage of Credible Intervals. A major advantage of using model-based methods to estimate disease location is that they can also provide a measure of the uncertainty of an estimate. To assess the accuracy of the estimated uncertainty for LATAG, we generated credible intervals of different sizes, ranging from 10% to 90%, based on the posterior distribution for each dataset. Figure 7 plots the number of datasets for which the disease mutation is located within each size credible interval and compares those numbers to the expected values. There is good accordance between the values, although it appears that for low and intermediate confidence levels, the constructed intervals are somewhat conservative and that the posterior distribution generated by LATAG slightly overestimates the uncertainty. The high uncertainty about the location of the disease mutation is reflected in the average size of the confidence intervals, which range from 0.06 cM for the 10%-CI to 0.85 cM for the 90%-CI.

For comparison, we also looked at the 95% confidence intervals that are generated by DHSMAP. Those intervals are considerably shorter than the intervals obtained from LATAG, at an average length of 0.37 cM for intervals obtained using their correction for pairwise correlation, and 0.15 cM without that correction. But for both models, the confidence intervals were too narrow, with 48% and 18%, respectively, of intervals containing the true disease locus (c.f., Morris et al., 2002).

In summary, LATAG seems to provide credible regions that are fairly well calibrated, or
Figure 8: *Comparison of the ability of LATAG (x-axis) and a SNP-based $\chi^2$ test (y-axis) to detect disease-causing loci in a test for association.* Each point corresponds to one of the 50 simulated datasets, and plots the most significant p-values obtained for that dataset using each method, corrected for multiple testing within the region. The dotted lines depict the $p=0.05$ cutoffs and the diagonal line plots the regression line through the log-log transformed data.

perhaps slightly conservative.

**Hypothesis testing.** To gauge the power of LATAG in a test for association, we assessed for each dataset whether we could detect the simulated region as a region harboring a disease mutation. To do this, we calculated the likelihood ratio at each focal point according to equation (1) and considered the maximal LR that we observed among all focal points as the evidence for association. For comparison, we also tested each SNP for association with the phenotype, using a standard Pearson $\chi^2$ test. We obtained an average maximum value of twice the log-likelihood ratio of 5.8 and an average maximum SNP-based $\chi^2$ of 8.5. In 88% of the simulations, the $\chi^2$-test generated a more significant singlepoint p-value.

However, because the extent of multiple-testing may be different for the two methods, this is not exactly the right comparison. The LATAG analysis consists of 50 tests, many of which are highly correlated, because the trees may differ little from one focal point to the next. The SNP-based test consists of about the same number of tests (one for each marker), and the correlation between tests depends on the LD between the markers. Therefore, a simple Bonferroni correction is too conservative for both test statistics. To perform tests that take the dependence structure in the data into account, we obtained p-values for each of the two test statistics by permutation (see “Simulation methods”). We observed that correcting for multiple tests has a strong impact on the signal of the single point analysis.
For 24% of the datasets, the single point analysis produced a region-wide p-value < 0.05, while in 30% of the datasets, LATAG produced a p-value < 0.05. Furthermore, the two tests do not always detect the same datasets: 1/3 of all the datasets that showed a significant single-point score did not have a significant signal with LATAG, while 45% of all disease loci that were detected with LATAG were not detected with the single-point analysis (Figure 8). Hence, although LATAG appears to be more powerful on average, there may be some value in performing SNP-based tests of association as well as that approach may detect some loci that would not be detected by LATAG.

Testing and Applications: Real data

To further illustrate our method we report analyses of two sets of case-control data. One dataset was used to map the gene responsible for cystic fibrosis, a simple recessive disorder (Kerem et al., 1989), while the other dataset is from a positional cloning study of a complex disease, Type 2 diabetes (Horikawa et al., 2000).

Example application 1: Cystic Fibrosis. The cystic fibrosis (CF) dataset used by Kerem et al. (1989) to map the CFTR locus has been used to evaluate several previous fine-mapping procedures, thus allowing an easy comparison between LATAG and other multipoint methods. The dataset was generated to find the gene responsible for cystic fibrosis (CF), a fully penetrant recessive disorder with an incidence of 1/2500 in Caucasians. Many different disease-causing mutations have been observed at the CFTR locus, but the most common mutation, ΔF508, is at quite high frequency, accounting for 66% of all mutant chromosomes.

The dataset consists of 23 RFLPs distributed over 1.8 Mb; these were genotyped in 47 affected individuals. In addition, 92 control haplotypes were obtained by sampling the non-transmitted parental chromosomes. High levels of association were observed for almost all markers in the region; the marker with the highest singlepoint association ($\chi^2 = 63$) is located at 870 kb from the left-hand end of the region. The ΔF508 mutation is at 885 kb, and is present in 62 of the 94 case chromosomes.

We ran ten independent runs of the Markov chain, estimating the average posterior likelihood at each of fifty evenly distributed points across the region. Each run had a burn-in of $2.5 \cdot 10^6$ steps for the first focal point and $10^6$ steps for each following focal point. In each run, we sampled 50 trees at each focal point, with a thinning interval of 10,000 steps. The runs took 8 hours each on a Pentium III processor. For each tree $T_i$, we calculated $P(\Phi|T_i, x)$ with the peeling algorithm. As the resulting posterior likelihoods seemed to be heavily dependent on a few outliers, we estimated the likelihood $P(x|\Phi, G)$ at each position $x$ by taking the median of the likelihoods $P(\Phi|T_i, x)$ instead of the average suggested by theory. As before, we used the posterior mode as our point estimate for location. Missing
Figure 9: Repeatability across runs: average posterior likelihoods for ten independent LATAG analyses of the CF dataset. For the location of the disease locus and the resulting posterior credible region refer to Figure 10.

data were imputed using PHASE 2.0 (Stephens et al., 2001).

**Results.** To provide a simple check of convergence, Figure 9 shows the results from the ten independent analyses of the CF dataset. As can be seen, all ten runs have modes in the same region and yield the same conclusion about the location of the causative variation.

Figure 10 summarizes our results across the ten runs. The posterior distribution is sharply peaked at 867 kb, near the true location of ΔF508 (which is at 885 kb). The 95% credible interval is rather narrow, extending from 814 kb to 920 kb. Even though there are several markers with little association to the trait in the vicinity of the deletion (Figure 10), the LATAG estimate is quite accurate. It is useful to compare our results to those obtained by other multi-point methods (see Table 1, modified from Morris et al. (2002)). For this dataset, most of the highest singlepoint $\chi^2$ values lie to the left of the true location of ΔF508, and so most of the methods err to the left, with some of the earliest methods (Terwilliger 95) actually excluding the true location from the confidence interval. Notice that the LATAG estimate is closer to the true location, and that the 95% credibility region is narrower than obtained by any of the previous methods.

To assess the ability of LATAG to detect the CF region by association, we calculated a likelihood ratio according to (1) and obtained $2 \cdot \ln(LR) = 40$. Assuming a $\chi^2$-distribution with one degree of freedom, this log likelihood ratio has an associated p-value of $3.7 \cdot 10^{-10}$. While this is extremely significant, it is less significant than obtained from simple tests of
Figure 10: Average posterior likelihoods generated by LATAG for the CF dataset (Kerem et al., 1989). The dots depict the association signals of the individual markers as $\chi^2$ statistics (see scale on right-hand side of figure). We display the likelihoods here because the posterior density is extremely peaked, with 95% of its mass inside the box marked by the dashed lines.

<table>
<thead>
<tr>
<th>Method</th>
<th>estimate</th>
<th>Variability</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terwilliger (1995)</td>
<td>770</td>
<td>690-870 [99.9% support interval]</td>
<td></td>
</tr>
<tr>
<td>McPeek and Strahs (1999)</td>
<td>950</td>
<td>440-1460 [95% confidence interval]</td>
<td>pairwise correction</td>
</tr>
<tr>
<td>Morris et al. (2000)</td>
<td>800</td>
<td>610-1070 [95% credible interval]</td>
<td>pairwise correction</td>
</tr>
<tr>
<td>Liu et al. (2001)</td>
<td>-</td>
<td>820-930 [95% credible interval]</td>
<td></td>
</tr>
<tr>
<td>Morris et al (2002)</td>
<td>850</td>
<td>650-1000 [95% credible interval]</td>
<td></td>
</tr>
<tr>
<td>LATAG</td>
<td>867</td>
<td>814-920 [95% credible interval]</td>
<td></td>
</tr>
</tbody>
</table>

Table 1: Estimates of locations of the CF-causing allele, as taken from Morris et al. (2002). The $\Delta F508$ mutation, which is responsible for 66% of all CF cases is located at position 885. Only estimates that are based on the entire dataset are presented.
association with individual SNPs (six of which yield $\chi^2$ values $> 50$). This may indicate that our likelihood ratio test does not fit the $\chi^2$ approximation very well, particularly far out in the tail of the distribution, or that our test is slightly less powerful in this extreme setting. Due to the extremely high level of significance, it is infeasible to generate an accurate p-value by permutation.

**Example application 2: Calpain-10.** Our second application comes from a positional cloning study that was searching for disease variation underlying Type 2 diabetes. Type 2 diabetes is the most common form of diabetes and in developed countries, it affects 10-20% of individuals over the age of 45 (Horikawa et al., 2000). This appears to be a highly complex disease, with no gene of major effect, and with environmental factors playing an important role. A linkage study in Mexican Americans localized a susceptibility gene to a region on chromosome 2 containing three genes, RNEPEPL1, CAPN10 and GPR35. A dataset that was generated by Horikawa et al. (2000) for positional cloning, consists of 85 SNPs distributed over an area of 876 kb. The markers were genotyped in 108 cases and 112 controls. No individual marker shows high association; the marker with the highest LD ($\chi^2 = 9$) is located at 121 kb from the left-hand end of the region. The original study also used some additional information from family sharing patterns that we do not consider here. Based on detailed analysis of those data, Horikawa et al. (2000) proposed that a combination of two haplotypes, each consisting of 3 SNPs within the CAPN10 gene, increases the risk of diabetes by 2- to 5-fold. The three SNPs that make up the haplotype are located at 121 kb, 124 kb and 134 kb.

We used the PHASE 2.0 algorithm, with recombination in the model (Stephens et al., 2001), to impute the phase information and missing genotypes for both cases and controls. Then we used LATAG to infer the posterior distribution of the location of the disease mutation and a p-value for association, as described above. Performing 8 independent runs of the MCMC, we generated a total of 800 draws from the posterior distribution $P(T|G, x)$ for each of 50 positions in the sequence. Each run had a burn-in of $5 \cdot 10^6$ steps for the first point and $10^6$ steps for each following point, a thinning interval of 10,000 steps between draws, and took 36 hours on a Pentium III processor.

**Results.** Figure 11 plots the estimated posterior distribution for the location of diabetes-associated variation in this region. From this distribution we estimate the position of the disease mutation at 131 kb, at the same location as the SNPs that Horikawa et al. (2000) reported as defining the key haplotypes. However the posterior distribution is quite wide, with 50% of its mass between 70 kb and 245 kb. The full 95% credibility region extends between 0 kb and 660 kb, indicating that we can only really exclude the right-hand end of this region. We would need larger samples to obtain more precision.

To assess whether we would have detected this region by association, based on this
Figure 11: Posterior distribution for the location of the diabetes-affecting variant(s) in the calpain-10 region (solid line; see scale on the left-hand side of the figure). The dots represent the association of individual markers with the phenotype (see $\chi^2$ scale on the right-hand side of the figure). The red dots indicate the three markers that define the disease-associated haplotypes reported by Horikawa et al. (2000).
dataset, we evaluated (1) and obtained $2 \ln(\text{LR}) = 6.0$ at the posterior mode, corresponding to a singlepoint p-value of $5.3 \cdot 10^{-4}$. When we correct for multiple testing using the simulation procedure, the overall significance level drops to a mildly significant $p=0.02$. In contrast, if we assess the significance of the highest observed singlepoint chi-square by permuting case and control labels, we obtain a nonsignificant p-value of 0.11.

Overall, our results are consistent with the conclusion of Horikawa et al. (2000), that CAPN10 is the gene that was responsible for their diabetes linkage signal in this region. However, our analysis cannot exclude GPR35 as the disease gene. As with the original analysis, our strongest signal is in the CAPN10 region, but our overall signal is only modestly significant.

**Discussion**

We have described a new unified method, LATAG, for association mapping and fine mapping with multipoint data. Our approach, based on a local approximation to the ARG, strikes a compromise between modeling the population genetic processes that produce the data, and the need for a model that is computationally tractable for large datasets.

Our association mapping method is similar in spirit to earlier tree-based methods (e.g., Templeton et al., 1987). However, we take a more probabilistic approach in the sense that we average over the uncertainty in trees, and consider an explicit mutation model at the unobserved disease locus. A more fundamental difference is that our tree inference scheme aims to model recombination explicitly, while the earlier methods make the most sense in small regions without evidence for recombination. Moreover, even for estimating the tree within a haplotype block, markers outside the block may contain additional information about that tree. It is typical for there to be at least some LD between haplotype blocks (Daly et al., 2001), and patterns of haplotype sharing across multiple blocks are potentially quite informative about the order of recent coalescent events. In our method, rather than forcing the user to predefine regions of limited recombination, the algorithm “adapts” to the data, in the sense that quite large regions of shared haplotypes may help to resolve recent coalescent events, while much smaller regions (e.g., corresponding to haplotype blocks), may be the relevant scale for reconstructing the topology of the more ancient coalescent events. Hence, we gather information both from mutation, and from recombination events to reconstruct the ancestral trees. By doing so, we can detect association even when there is allelic heterogeneity, and we can gain information about low frequency disease mutations, even using only intermediate frequency SNPs. Our simulations indicated that LATAG is substantially more powerful than singlepoint SNP-based tests of association, at least for the scenario considered.

It is also natural to compare LATAG to recent fine mapping algorithms. One major difference between LATAG and most of the previous coalescent-based algorithms is that
LATAG aims to reconstruct the ancestry of all the sampled chromosomes, not just of case chromosomes. By considering the ancestry of all individuals at once, we can deal with more general phenotype models and we can model allelic heterogeneity and incomplete penetrance in a natural way, although including this additional information may increase computation time. This also represents the first multipoint fine mapping method for quantitative traits. Our approach can also produce penetrance estimates under haploid and diploid models (the latter to be presented elsewhere), though these estimates may not be straightforward to interpret when the ascertainment of samples is not random. We have not focused on this here, but our approach also produces a posterior probability that each chromosome carries a disease mutation. This can be used to guide full resequencing of implicated regions. Including control chromosomes in the tree allows us to make better use of the control data than earlier methods that just used controls to estimate the SNP allele frequencies; as exemplified by the strong performance of our method on the CF data and on the simulated data.

For any model-based approach such as LATAG, it is worth considering the various modeling assumptions, and how these might affect the results. In general, it seems that most of the inaccuracies of the model can be overcome by informative data; at worst they might slightly reduce our power and precision (c.f., Morris et al., 2002). For instance, the Markov model that we use for LD outside the inherited region is not strictly accurate, and might be expected to produce a slight bias toward keeping too much sequence on the tree. Inaccuracies there may explain the tendency toward conservativeness in the coverage of our credible intervals (Figure 7). Similarly, the finite sites model used for SNP mutation, ignoring SNP ascertainment, is clearly inaccurate. However, this model is computationally convenient, and it seems likely that the data should overwhelm deficiencies here; again, that view is supported by the results. Besides, allowing recurrent SNP mutation provides an ad hoc way of allowing for gene conversion, which might otherwise confound our inference.

Another issue is that we ignore the ascertainment of cases, and natural selection on the disease variants. These processes are expected to distort the shape of the tree (cases will be more closely related than predicted under the coalescent model). But this distortion effect will be most pronounced when the signal is very strong (e.g., for a recent highly penetrant mutation, as in the CF data). In that case, the data will usually be strong enough to overwhelm the rather weak coalescent prior. In other words, when the coalescent model is furthest from the truth, the data are likely to be very informative and should override the misspecifications of the prior.

At present we also assume that even if there are multiple disease mutations in a region, that all of these occur at essentially the same position. This may well be a poor assumption. For example, mutations in different exons of a single gene might be many kilobases apart. Neither our method, nor any other existing method would handle this well; however dealing with this issue is surely an important problem for the future.

Full-likelihood coalescent methods such as LATAG pose considerable computational chal-
It is not easy to design MCMC algorithms that can traverse through tree space efficiently and produce robust, repeatable results for large data sets (Wall, 2000). As noted above, we use a local approximation to the ARG to substantially simplify the space that we are mixing through. We also clip off upper parts of the sequence that are not inherited (Figures 3, 4) to further improve mixing. In our MCMC design, we chose to augment the data by storing, and mixing over, the sequence at internal nodes. Doing so makes the Metropolis-Hastings calculations for each proposed update extremely fast and allows the algorithm to use the sequence identity to propose more effective tree rearrangements. On the downside, storing this extra information may plausibly impede mixing; further experimentation will be required to help determine the best design. The calpain-10 dataset, consisting of 85 SNPs genotyped in 440 chromosomes, is at the upper end of what our current algorithm can handle reliably (and is also very large by the current standards of other full-likelihood coalescent methods). One advantage of our approach over other genealogy-based methods, is that LATAG can easily be parallelized, since the trees are reconstructed independently of each other at different focal points and the analysis of trees can be performed independently of their generation process. Each of these operations can run on a different processor and LATAG can make efficient use of modern computing facilities. Nevertheless, improving the algorithm to deal with larger datasets is a focus of our ongoing research.

One question that arises in this context is whether to treat haplotype phase from diploid genotypes as known when inferring the trees (here we estimated phase using *Phase* 2.0 Stephens et al., 2001). The alternative—which is more statistically sound—is to use the MCMC coalescent algorithm to mix over unknown phase along with tree topology. Morris et al. (2004) implemented such a method, and reported that it produced more accurate fine mapping results than did a method that used haplotypes estimated by a simple EM algorithm. However, given that it is already difficult to achieve good MCMC performance in large datasets of known haplotypes, it is unclear to us that also mixing over haplotypes is necessarily a good strategy. We look forward to further research on this issue.

In summary, our new methods provide a coherent framework for achieving different goals of LD-based mapping. Furthermore, they perform well on real and simulated data, compared to standard existing methods. One of the biggest challenges for the future will be to develop our current framework so that it can be applied to the massive datasets that will soon be forthcoming in the human genetics community.

**Acknowledgments**

We thank Nancy Cox, Dick Hudson, Dan Nicolae, Matthew Stephens and two anonymous reviewers, plus the other members of our lab, for helpful discussions and/or comments on the manuscript, and Nancy Cox for giving us access to the calpain-10 data. This work was supported by grant HG 2772 to JKP from the National Institutes of Health.
References


Appendix 1: The MCMC algorithm

The goal of the Markov chain Monte Carlo algorithm is to generate trees from the distribution \( Pr(T_x | X = x, G) \). As before, \( G \) denotes haplotype data across some region, and \( x \) is a focal point within that region. We treat the unknown genealogy, node times, and ancestral sequence at each node as missing data, and use MCMC to integrate over these missing data. For a more detailed description, we need to introduce some more notation. As an aid to the reader, the notation used in this appendix is summarized in table 2.

We assume that recombination events and mutation events on one side of \( x \) occur independently of events on the other side of \( x \), so that conditional on the tree topology and branch lengths, the full likelihood can be computed as the product of the left-hand and right-hand likelihoods. Thus, it is sufficient to describe mutations and recombinations on the right side of \( x \) with the understanding that the same process occurs on the left side. Let us therefore assume, without loss of generality, that \( x \) is immediately to the left of marker 1.

The sample we are looking at consists of \( n \) chromosomes that are typed at \( L \) loci. The marker map can be described by the following variables. Let \( d_i, i \in \{1, \ldots, L\} \) be the physical distance between the focal point \( x \) and marker \( i \) and let \( \alpha_i \) be the number of alleles at marker \( i \). Let \( \theta/2 \) be the mutation rate of each marker per unit coalescent time, and let \( \rho/2 \) be the recombination rate per unit coalescent time per unit distance. That is, recombination events occur between the focal point \( x \) and marker \( i \) at rate \( \rho \cdot d_i/2 \) per unit coalescent time.

The coalescent tree \( T_x \) is described by the following two variables: Let \( \Omega = (\omega_n, \ldots, \omega_2) \) denote the times between successive coalescent events (e.g., \( \omega_n \) is the time during which there are \( n \) lineages in the tree). Let \( \tau \) denote the labeled topology of the tree. For notational purposes, it is convenient to introduce \( K_j, j \in \{1, \ldots, 2n-1\} \) as the ordered set of nodes on the tree, so that \( \{K_1, \ldots, K_n\} \) are the external nodes, \( K_{n+1} \) is the node of the first coalescent event and \( K_{2n-1} \) is the MRCA. Furthermore, let \( B = (b_1, \ldots, b_{2n-2}) \) be the vector of branch lengths, where \( b_j \) is the branch length between node \( K_j \) and its parental node.

Let \( s_i^j \in \{1, \ldots, \alpha_i\} \) be the observed or inferred allele at node \( K_j \) at marker \( i \), and \( s^j \) denote the full haplotype \( \{s_1^j, s_2^j \ldots s_L^j\} \). If marker \( i \) has no sequence in \( K_j \), then \( s_i^j \) is set to 0. Let \( S = (s^1, \ldots, s^{2n-1}) \) be the vector of all sequence information in internal and external nodes. In this notation \( G = \{s^1, \ldots, s^n\} \). Let \( r_j \in \{1, \ldots, L+1\} \) be the marker closest to \( x \) that is not inherited to node \( K_j \) from its ancestor due to recombination, where \( r_j = L+1 \) indicates that the entire set of markers is inherited. Let \( R \) be the vector of all \( r_j \). Using this notation, the goal of our algorithm is to sample from \( Pr(\Omega, \tau | s^1, \ldots, s^n) \), while treating \((\theta, \rho, R, \ s^{n+1}, \ldots, s^{2n-1}) \) as augmented data. Let \( A = (\Omega, \tau, \theta, \rho, R, \ s^{n+1}, \ldots, s^{2n-1}) \) be the joint vector of unknown parameters. At each step of the algorithm, we draw a candidate value \( \tilde{A} \) from a proposal density \( O(\cdot | A, G) \). Details about the proposals are given below. The candidate value \( \tilde{A} \) is accepted to replace \( A \) with probability \( \alpha(A, \tilde{A}) \), where

\[
\alpha(A, \tilde{A}) = \min\{1, \frac{Pr(\tilde{A} | G) \cdot O(A | \tilde{A}, G)}{Pr(A | G) \cdot O(A | A, G)}\}
\] (10)
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x$</td>
<td>focal point</td>
</tr>
<tr>
<td>$T_x$</td>
<td>bifurcating tree at locus $x$</td>
</tr>
<tr>
<td>$G$</td>
<td>genotypes (marker data)</td>
</tr>
<tr>
<td>$n$</td>
<td>number of chromosomes in the sample</td>
</tr>
<tr>
<td>$L$</td>
<td>number of markers</td>
</tr>
<tr>
<td>$d_i$</td>
<td>distance between $x$ and marker $i \in {1, \ldots, L}$</td>
</tr>
<tr>
<td>$\alpha_i$</td>
<td>number of alleles at marker $i$</td>
</tr>
<tr>
<td>$\theta / 2$</td>
<td>mutation rate of each marker per coalescent time unit</td>
</tr>
<tr>
<td>$\rho / 2$</td>
<td>recombination rate per kb per coalescent time unit</td>
</tr>
<tr>
<td>$\omega_i$</td>
<td>times between coalescent events</td>
</tr>
<tr>
<td>$\Omega$</td>
<td>vector of coalescent times $\omega_i$</td>
</tr>
<tr>
<td>$\tau$</td>
<td>labeled topology of the tree</td>
</tr>
<tr>
<td>$K_j$</td>
<td>ordered set of nodes on the tree</td>
</tr>
<tr>
<td>$b_j$</td>
<td>branch length between node $K_j$ and its parental node</td>
</tr>
<tr>
<td>$B$</td>
<td>vector of branch lengths $b_j$</td>
</tr>
<tr>
<td>$s^j_i$</td>
<td>sequence information in node $K_j$ at marker $i$</td>
</tr>
<tr>
<td>$r_j$</td>
<td>marker closest to $x$ that is not inherited to node $K_j$ due to recombination</td>
</tr>
<tr>
<td>$R$</td>
<td>vector of recombinations $r_j$</td>
</tr>
<tr>
<td>$A$</td>
<td>$(\Omega, \tau, \theta, \rho, R, s^{n+1}, \ldots, s^{2n-1})$, the joint vector of unknown parameters.</td>
</tr>
</tbody>
</table>

Table 2: Summary list of notation for the MCMC. For a more detailed description refer to the text.
is the usual Metropolis-Hastings Ratio; otherwise the old value \( A \) is retained. The probabilities \( Pr(\tilde{A} \mid G) \) and \( Pr(A \mid G) \) are calculated according to the details given below. As is standard, the initial steps of the Markov chain are discarded, as they are heavily influenced by the starting condition. Inference is then performed on the subsequent set of topologies, with an appropriate thinning interval.

To evaluate (10), we need to calculate the probabilities \( Pr(\tilde{A} \mid G) \) and \( Pr(A \mid G) \). To this end, we first need to establish some basic models for recombination and mutation.

**Mutation Model.** We assume a finite sites mutation model with parent-independent mutation at rate \( \theta/2 \) per branch, per unit coalescent time. That is, at each marker, mutations occur as a Poisson process at rate \( \theta/2 \), and the new allele following a mutation is drawn uniformly at random from the \( \alpha_i \) possible alleles. (Hence, at a site with 2 alleles, \( \theta/2 \) is twice the biological mutation rate, which only counts mutations that change the allele.) It should be pointed out, that \( \theta \) as it is used here represents the mutation rate of a pre-ascertained SNP, not the usual mutation rate of a random base pair. Let node \( K_l \) be ancestral to \( K_k \), then conditional on the fact that no recombination occurs between \( x \) and marker \( i \) between \( K_l \) and \( K_k \), the allelic state of marker \( i \) has the following distribution

\[
Pr(s_i^k = a_1 \mid s_i^l = a_2, b_k, r_k > i) = \begin{cases} 
\frac{1}{\alpha_i} \cdot (1 - e^{-\theta b_k/2}) & \text{if } a_1 \neq a_2 \\
\frac{1}{\alpha_i} \cdot (1 - e^{-\theta b_k/2}) + e^{-\theta b_k/2} & \text{if } a_1 = a_2 
\end{cases}
\]

(11)

where \( a_1, a_2 \in \{1, \ldots, \alpha_i\} \).

**Background Haplotype Probabilities.** For the model of the recombination process it is necessary to provide the probability that a haplotype could arise on the part of the ancestry that is not described by \( T_x \). Let \( q, v \in \{1, \ldots, L + 1\} \) be positions on the marker map and \( (s_i)_{q \leq i < v} \) the sequence between those two positions. Then \( H((s_i)_{q \leq i < v}) \) designates the probability of drawing the haplotype \( (s_i)_{q \leq i < v} \) from the population. As in some previous work in this area (e.g., McPeek and Strahs, 1999; Morris et al., 2002), we model the likelihood for sequence that recombines into the tree as a first-order Markov process, estimating the allele frequencies and two-site haplotype frequencies as proportional to the sample frequencies plus 1.

**Recombination Model.** Let the nodes \( K_k, K_q \) being the descendants of node \( K_j \), then \( z_j \in \{1, \ldots, L + 1\} \) is defined as \( z_j = \max\{r_k, r_q\} \). Thus \( z_j \) is the marker closest to \( x \) in node \( K_j \) that will not be inherited to the present. For \( i \in \{1, \ldots, n\} \), by definition \( z_i = L + 1 \). Then the probabilities of recombination events on the branch from \( K_j \), conditional on the
state at $K_j$ are

$$\Pr(r_j = c \mid b_j, z_j) = \begin{cases} 
0 & \text{if } c > z_j \\
\int_0^{d_c} b_j \rho / (2e^{-b_j\rho t/2}) dt & \text{if } c = 1 \\
\int_{d_c}^{d_{c-1}} b_j \rho / (2e^{-b_j\rho t/2}) dt & \text{if } 1 < c < z_j \\
\int_{d_{c-1}}^{\infty} b_j \rho / (2e^{-b_j\rho t/2}) dt & \text{if } c = z_j 
\end{cases}$$  \hspace{1cm} (12)$$

for $c \in \{1, \ldots, L + 1\}$. For some rearrangements in the tree in the MCMC, it is necessary to calculate the probability of a recombination on $b_j$, conditional on the state of nodes that are ancestral to $K_j$. In our model, the amount of information at a node is dependent on the recombination events that occur on branches descending from that node. Thus certain rearrangements may be incompatible with the rest of the tree, as they may provide an upper bound to sequence lengths in ancestral nodes. If the sequence in an ancestral node is longer than this upper bound, the resulting tree impossible. This has to be taken into account when calculating the probability of a given recombination conditional on the sequence length at nodes. Let $K_u$ be a node that is ancestral to $K_j$. Then, $r_j$ is consistent with $z_u$ if it allows sequence information up to $z_u$ to reach node $K_u$. Then, the distribution for recombination events in $K_j$ is

$$\Pr(r_j = c \mid b_j, z_j, z_u) \propto \begin{cases} 
0 & \text{if } c \text{ not consistent with } z_u \\
\Pr(r_j = c \mid b_j, z_j) & \text{if } c \text{ consistent with } z_u 
\end{cases}$$

Prior probabilities. In our current implementation, the priors for $\Omega$ and $\tau$ are those given by the standard neutral model for a single locus. That is, $\omega$, the time during which there are $l$ lineages, is exponentially distributed with parameter $\frac{1}{2}$, independently for each $l$. The topology $\tau$ is a bifurcating tree with $n$ labeled tips; when there are $l$ lineages, the probability that two particular lineages coalesce is $\frac{1}{2}$ for all pairs. The priors for $\theta$ and $\rho$ are taken as uniform.

Probability of a tree. With these models in place, we can now write the probability of the tree and augmented data, conditional on the observed data, as

$$\Pr(\Omega, \tau, \theta, \rho, R, s^{n+1}, \ldots, s^{2n-1} \mid s^1, \ldots, s^n) = \frac{\Pr(\Omega, \tau) \cdot \Pr(\rho) \cdot \Pr(\theta)}{\Pr(s^1, \ldots, s^n)} \cdot \Pr(R, S \mid \Omega, \tau, \rho, \theta),$$

assuming independence of the prior probabilities for $(\Omega, \tau, \rho, \theta$. The prior probabilities in (13) (i.e., $\Pr(\Omega, \tau)$, $\Pr(\rho)$ and $\Pr(\theta)$ are computed as above. $\Pr(s^1, \ldots, s^n)$ is constant and cancels out of the Metropolis-Hastings ratio. The last factor can be calculated as

$$\Pr(R, S \mid \Omega, \tau, \rho, \theta) = \Pr(R \mid \Omega, \tau, \rho) \cdot \Pr(S \mid R, \Omega, \tau, \rho)$$

(14)

Now, as the nodes are ordered by their time since the present, we can calculate the first term of (14) as

$$\Pr(R \mid \Omega, \tau, \rho) = \Pr(r_1 \mid \Omega, \tau, \rho) \cdot \Pr(r_2 \mid r_1, \Omega, \tau, \rho) \cdot \ldots \cdot \Pr(r_{2n-2} \mid r_1, \ldots, r_{2n-3}, \Omega, \tau, \rho).$$

(15)
The individual terms in (15) can be calculated according to (12), as for any node $K_j$ the recombinations for all nodes "below" $K_j$ are in the conditional, therefore $z_j$ is known. The second term of (14) can be calculated as

$$\Pr(S \mid R, \Omega, \tau, \theta) = \Pr(s^{2n-1} \mid R, \Omega, \tau, \theta) \cdot \cdots \cdot \Pr(s^1 \mid s^2, \ldots, s^{2n-1}, R, \Omega, \tau, \theta)$$  \hspace{1cm} (16)$$

The first term of (16) represents the sequence at the MRCA of the coalescent tree and can be approximated by drawing from $H((s_i^{2n-1})_{y_i \leq z < z_{2n-1}})$. Every other term in (16) calculates the probability of sequence in node $K_v$ conditional on the sequence of its ancestral node $K_j$ while nodes that are descendants of $K_v$ are not in the conditional. Therefore, it can be written as

$$\Pr(s^v \mid s^j, b_v, z_v, r_v, \tau) = H((s_i^{2n-1})_{r_i \leq z < z_v}) \cdot \prod_{i=y_v}^{r_v-1} \Pr(s^v_i \mid s^j_i, b_v)$$

where the probabilities in the second term are calculated according to (11).

**MCMC updates.** The MCMC algorithm draws trees from $T \mid G$, while treating $(\theta, \rho, R, s^{n+1}, \ldots, s^{2n-1})$ as augmented data. We start with an initial value for each of these variables, chosen either at random from the prior, or using some heuristic guess. Then at each step of the algorithm, we propose a change of one or more parameters. Each step includes the "local update of internal nodes" for all nodes, and one or more of the topology rearrangements. The updates for $\theta$ and $\rho$ are performed less often. Each proposal is accepted according to the Metropolis-Hastings ratio (10). In the following, we describe the different proposals employed. Dependent on the nature of the dataset, we perform the different proposal at different rates. For every parameter $z$ let $\tilde{z}$ denote the proposed new parameter. Furthermore, we define $t_i$ to be the time between node $K_i$ and the present. The different changes we propose are:

**Propose new $\theta$.** A new $\tilde{\theta}$ is drawn from a uniform distribution on the interval $(0.5 \cdot \theta, 2 \cdot \theta)$.

**Propose new $\rho$.** A new $\tilde{\rho}$ is drawn from a uniform distribution on the interval $(0.5 \cdot \rho, 2 \cdot \rho)$.

**Local update of internal nodes.** Starting at the terminal nodes, we propose for each node $K_i$ an $\tilde{r}_i$, a time $\tilde{t}_i$ and a sequence $\tilde{s}^i$ conditional on the sequence and recombination events at surrounding loci. All nodes are visited in each step of the Markov chain.

**Major rearrangements.** We randomly select a node $K_i$ that can be removed from its location, without causing inconsistencies among its parental nodes (Figure 12). Using the notation illustrated in the Figure, $K_i$ is a candidate to be moved if

$$\max\{r_q, b_j\} \geq b_o.$$
Then we consider all other nodes whose parental nodes are older than $K_i$ and weight them according to their sequence similarity with $K_i$. Given those weights, we draw one node $K_c$. Let $K_k$ be the parental node of $K_c$. We then draw a time $t$ uniformly from the interval $(\text{max}\{t_c, t_i, t_k\})$ and propose a new tree where $K_i$ coalesces with $K_j$ at node $K_p$ time $t$, while $K_j$ and $K_q$ coalesce at node $K_o$ (see Figure 12). We draw new recombinations $\tilde{r}_j$, $\tilde{r}_c$, $\tilde{r}_i$ and $\tilde{r}_p$ and a new sequence $\tilde{s}^r$ conditional on the information at surrounding nodes.

**Minor rearrangements.** We draw an internal node $K_i$ of the tree. Let $K_i$ and $K_j$ coalesce at $K_p$ and $K_p$ and $K_q$ coalesce at $K_o$. Then we propose a tree, where $K_j$ and $K_q$ coalesce at $K_p$, and $K_i$ and $K_p$ coalesce at $K_o$, while the coalescent times remain unchanged. We also propose a new $\tilde{s}^p$ and $\tilde{r}_p$.

**Reordering of coalescent events.** We select a internal node $K_i$ that has the direct descendants $K_k$ and $K_i$ and the parental node $K_m$. Then we select a second internal node $K_j$ that has the direct descendants $K_o$ and $K_p$ and the parental node $K_q$ with $t_j \in (\text{max}\{t_k, t_l, t_m\})$ and $t_i \in (\text{max}\{t_o, t_p, t_q\})$ and propose an exchange of times $t_i$ and $t_j$.

**Appendix 2: Calculating the phenotype likelihoods**

Given a tree $T_x$, we need to compute $Pr(\Phi|X = x, T_x)$, the probability of observing the arrangement of phenotypes on the tree. To do this, we assume that all disease mutations occur as a Poisson process with rate $\nu/2$. Furthermore, we assume that multiple mutations on the same chromosomes have no further effect, thus every chromosome that carries at least one mutation has the same distribution of phenotypes. Under this model we have developed the following approach to calculate $Pr(\Phi|X = x, T_x)$. 

---

Figure 12: Proposal of major rearrangements of the topology of tree. Nodes $K_i$ are displayed as $i$. One node ($I$) and all its descendants are moved from one clade of the tree to a different clade.
**Peeling algorithm.** Recall that $P^\phi_m$, denotes the probability that a chromosome comes from an individual with phenotype $\phi$ given that it has mutation ($m = 0$) or at least one mutation ($m = 1$). Then $Pr(\Phi|X = x, T_x)$ can be calculated exactly, using the peeling algorithm (Felsenstein, 1981).

Let $m_i$ be an indicator for the mutation status at node $K_i$, where $m_i = 1$, if node $K_i$ carries at least one disease mutation and $m_i = 0$ otherwise. Furthermore, let $\Phi_i$ be the phenotypes of all terminal nodes that descend from node $K_i$. Then it is straightforward to calculate $Pr(\Phi_i|m_i = 1)$, as further mutations on branches below $K_i$ do not affect the phenotype (by assumption). Therefore,

$$Pr(\Phi_i|m_i = 1) = \prod_{K_j \text{ is terminal descendants of } K_i} P^\phi_{ij}$$

where $K_j$ denotes the terminal descendants of $K_i$. In the case of a case control-phenotype, where $\Phi_i$ consists of $a$ affecteds and $u$ unaffecteds, equation (17) can be written as

$$Pr(\Phi_i|m_i = 1) = (P^0_1)^u \cdot (P^1_1)^a,$$

where $u$ is the number of controls and $a$ is the number of cases among the terminal descendants of $K_i$.

On the other hand, $Pr(\Phi_i|m_i = 0)$ is a little more complicated to calculate. Here we can make use of the assumption that conditional on the mutation status phenotypes at each terminal node occur independently. Then mutations on branches are only affecting phenotypes that descend from this branch. Let $K_y$ be an internal node of the tree and $K_s, K_t$ be the descendants of $K_y$. Furthermore, let $\mu_i$ be the probability that there is at least one disease mutation on the branch from node $i$ to its parental node, calculated as $\mu_i = 1 - e^{-\nu b_i/2}$. Then we can write:

$$Pr(\Phi_y|m_y = 0) = (Pr(\Phi_s|m_s = 0) \cdot (1 - \mu_s) + Pr(\Phi_s|m_s = 1) \cdot \mu_s) \cdot (Pr(\Phi_t|m_t = 0) \cdot (1 - \mu_t) + Pr(\Phi_t|m_t = 1) \cdot \mu_t)$$

While $Pr(\Phi_s|m_s = 1)$ and $Pr(\Phi_t|m_t = 1)$ can be calculated according to (17), $Pr(\Phi_s|m_s = 0) = P^\phi_0$, if $s$ is a terminal node. Therefore, we can calculate $Pr(\Phi_y|m_y = 0)$ for every internal node by starting at the most recent nodes and working iteratively backwards in time. As $Pr(\Phi|X = x, T_x) = Pr(\Phi_{MRCA}|m_{MRCA} = 0)$, this allows us to calculate the likelihood of the phenotypes given the tree.

**Integrating over penetrances.** The calculations just described are for fixed values of $P^\phi_0$ and $P^\phi_1$. Since these are unknown in advance, our Bayesian computations are based on integrating over the space of possible penetrances. To this end, let us assume, these probabilities are governed by a vector $P$ of variables that live on the bounded set $\Delta$. In the
case of a binary phenotype, this vector $P$ consists of the penetrances of the carriers/non-carriers and $\Delta = [0, 1] \times [0, 1]$, while for a normally distributed quantitative phenotype it is composed of the variances and the means of carriers and non-carriers. Then we want to evaluate

$$Pr(\Phi|T_x, X = x) = \int_{P \in \Delta} Pr(\Phi|T_x, X = x, P) Pr(P) dP.$$  

In practice, we are unable to calculate the integral. We therefore substitute

$$Pr(\Phi|T_x, X = x) \approx \frac{1}{k} \sum_{i=1}^{k} Pr(\Phi|T_x, X = x, P_i)$$

with $\hat{\Delta} = \{P_1, \ldots P_k\}$ selected from a suitable grid on $\Delta$. For the analysis presented here, we used $(P_0^0, P_1^0) \in \hat{\Delta} = \{(0.05, 0.05), (0.1, 0.05), \ldots, (0.95, 0.95)\}$.