Bayesian Population Genomic Inference of Crossing Over and Gene Conversion

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ABSTRACT
Meiotic recombination is a fundamental cellular mechanism in sexually reproducing organisms and its different forms, crossing over and gene conversion both play an important role in shaping genetic variation in populations. Here, we describe a coalescent-based full-likelihood Markov chain Monte Carlo (MCMC) method for jointly estimating the crossing-over, gene-conversion, and mean tract length parameters from population genomic data under a Bayesian framework. Although computationally more expensive than methods that use approximate likelihoods, the relative efficiency of our method is expected to be optimal in theory. Furthermore, it is also possible to obtain a posterior sample of genealogies for the data using this method. We first check the performance of the new method on simulated data and verify its correctness. We also extend the method for inference under models with variable gene-conversion and crossing-over rates and demonstrate its ability to identify recombination hotspots. Then, we apply the method to two empirical data sets that were sequenced in the telomeric regions of the X chromosome of Drosophila melanogaster. Our results indicate that gene conversion occurs more frequently than crossing over in the su-w and su-s gene sequences while the local rates of crossing over as inferred by our program are not low. The mean tract lengths for gene-conversion events are estimated to be ~70 bp and 430 bp, respectively, for these data sets. Finally, we discuss ideas and optimizations for reducing the execution time of our algorithm.

CROSSING over and gene conversion refer to two different mechanisms of recombination by which homologous chromosomes exchange genetic material during meiosis. In many eukaryotes, recombination is an essential requirement for sexual reproduction because it maintains physical connections between homologous chromosomes and contributes to proper segregation at the end of the first meiotic division. As currently understood, recombination starts with the formation of a double-stranded break in the DNA and proceeds by a series of biochemical steps leading to its repair (e.g., see Szostak et al. 1983; Stahl 1994). This repair can result in either the reciprocal exchange of large chromosomal regions (called crossing over) or the exchange of short DNA tracts (called gene conversion). The stretch of DNA exchanged during a gene-conversion event is called a conversion tract. A crossing-over event involves a single breakpoint in the chromosome and regions beyond this breakpoint are swapped between homologs to create mosaic products. In contrast, a gene conversion creates a mosaic in which a short internal fragment comes from one homolog while the rest of the chromosome flanking this fragment comes from the other. The products are similar in length to the participating homologs.

Crossing over is the better studied mechanism of recombination and crossing-over rates are known to vary tremendously across the genome at all scales. In particular, sperm-typing experiments as well as population genetic analysis of human polymorphism data have provided considerable evidence for fine-scale rate variation along the genome (e.g., see Fullerton et al. 1994; Dunham et al. 1999; Jeffreys et al. 2001, 2005; Innan et al. 2003; Crawford et al. 2004; McVean et al. 2004; International Hapmap Consortium 2005; Myers et al. 2005; Fearnhead and Smith 2005; Tiemann Boege et al. 2006; Coop et al. 2008). Notably, the population genetic studies ignored the effects of gene conversion and assumed that crossing over was the only mechanism underlying all exchanges. In general, gene conversion has been harder to study due to lack of appropriate fine-scale data and powerful statistical tools. Sperm-typing studies at a few individual human loci have provided
evidence of high rates of gene conversion relative to crossing over (Zangenberg et al. 1995; Jeffreys and May 2004) and observed tract lengths that appear to be in the range of 50–150 bp. Gene conversion has also been studied experimentally in yeast and fruit flies (Fogel et al. 1983; Hilliker et al. 1991, 1994; Paques and Haber 1999; Allers and Lichten 2001; Mancera et al. 2008) and tracts are estimated to be in the range of 350–2000 bp in these organisms. Nevertheless, the nature and extent of variation in gene-conversion rates along genomes remains by and large unknown.

Characterizing the basic parameters of gene conversion (i.e., rates and tract lengths) within organisms is useful for a variety of reasons. First, it helps to better explain the patterns of linkage disequilibrium observed in single-nucleotide polymorphism (SNP) data (Wall and Pritchard 2003; Padhukasahasram et al. 2004). Second, algorithms for association mapping (e.g., Liu et al. 2001; Morris et al. 2002; Niu et al. 2002; Carlson et al. 2004) and for detecting natural selection (e.g., Voight et al. 2006) make assumptions about the local recombination parameters and conversion estimates may help fine-tune such methods. Finally, meiotic gene conversion is a fundamental biological mechanism and its relationship to crossing over is not yet fully understood and is an important open question. The emergence of dense SNP data sets from next generation sequencing technologies in the coming years presents a major opportunity to accurately quantify recombination rates from a population genetics standpoint. Thus, population genetic analysis of gene-conversion and crossing-over rates can eventually shed light on the relationship between these two processes and can supplement experimental approaches in answering this question (see previous studies Langley et al. 2000; Allers and Lichten 2001; Andolfatto and Wall 2003; Padhukasahasram et al. 2006; Mancera et al. 2008).

Currently, several statistical methods exist that are designed to jointly estimate the crossing-over and gene-conversion rates from population genetic data. Methods developed by Frisse et al. (2001), Ptak et al. (2004), and Wall (2004) generalize the composite-likelihood approach first proposed in Hudson (2001). These approaches divide the data into small subsets (pairs or triplets of segregating sites), calculate likelihoods for these subsets, and multiply them together. The likelihood thus obtained is called the composite likelihood. Composite-likelihood methods use precomputed likelihood lookup tables for all the possible configurations of the subsets and are typically fast. Because the subsets are not independent of one another, they do not calculate the true likelihood of the data. Therefore, correct confidence intervals can be obtained only by using simulations.

Padhukasahasram et al. (2006) describe a rejection-sampling method that simultaneously utilizes informative long-range and short-range summary statistics to infer the recombination parameters. This approach uses only part of the information available in the data for the sake of computational efficiency. Confidence intervals may be directly calculated from the likelihood surface in this method.

The approximate-likelihood method for estimating crossing-over rates [called product of approximate conditionals (PAC)] that was proposed in Li and Stephens (2003) has also been extended by several recent studies to include gene conversion. Briefly, this method infers recombination parameters under a heuristic model and is computationally efficient. However, inference is currently restricted to the constant population size Wright–Fisher model only. Hellenthal (2006) used a PAC model where the conversion tract can include at most one marker. Gay et al. (2007) improved on that work to allow for arbitrary gene-conversion tract lengths and this method can be used for coestimating crossing-over and gene-conversion rates as well as tract lengths. One simplification in their model was that they disallowed overlapping gene-conversion events. Yin et al. (2009) further generalized this work to allow for overlapping events and in theory this method is expected to perform at least as well as the method of Gay et al. (2007). Simulations indicate that this generalization (at least for a subset of parameters) leads to a more accurate PAC-based method for jointly estimating all three recombination parameters.

In this article, we extend the Bayesian Markov chain Monte Carlo (MCMC) method (originally developed to infer crossing over exclusively) of Wang and Rannala (2008) to jointly estimate the population crossing-over rate, the population gene-conversion rate, and the mean conversion tract length from SNP data. We first check the performance of the new method on simulated data and verify its correctness. In addition, we extend the method for inference under models with variable gene-conversion and crossing-over rates and demonstrate its ability to identify recombination hotspots. Next, we apply the method to two empirical data sets that were sequenced in the telomeric regions of the X chromosome of Drosophila melanogaster. Because the new method generates a posterior sample of genealogies consistent with the observed data, it is possible to obtain the distribution of gene-conversion breakpoints for a data set. Comparison of this distribution with a prior using a Bayes factor may be informative about the locations of conversion breakpoints in the history of the sample. We also calculate the Bayes factors for gene-conversion initiation points for these two Drosophila data sets. Our analysis suggests that gene conversion occurs more frequently than crossing over in the su-w and su-s gene sequences while the local rates of crossing over as inferred by our program are not low. The estimated mean tract lengths for these regions are ~70 bp and 430 bp, respectively. Furthermore, plots of the logarithm of Bayes factors for conversion start points do not indicate any strong deviations from a prior distribution expected under a model with uniform recombination rates. Finally, we discuss ideas and optimizations for improving the run-time efficiency of our algorithm.

Materials and Methods

Our recombination inference method is based on the retrospective coalescent framework in which the genealogy
of a sample of sequences is approximated by a graph called
the ancestral recombination graph (ARG) (Kingman 1982;
Hudson 1983; Griffiths and Marjoram 1996). In particular,
the method uses the coalescent with gene conversion as
described in Wiuf and Hein (2000). In this model, conver-
sion tract lengths are assumed to be geometrically distrib-
uted (Hilliker et al. 1994). The distribution of tract length
(Z) given that a gene-conversion event has occurred, i.e.,
P(Z = z|gene conversion), is thus equal to q(1 − q)z−1,
where q denotes the reciprocal of the mean conversion tract
length (m). Consider a sequence of length L + 1 bp. A gene-
conversion event can initiate between any two adjacent base
pairs along the sequence. Without loss of generality, the
model also assumes that a tract will always be to the right
of the initiation position. Thus, if a conversion event initiates
at gap S, S = 1, 2, 3, . . . , L, then the end point is S + z.
Gene-conversion events that initiate outside the sequence on
the left and terminate within the sequence must also be in-
cluded in our analysis. The probability that the tract length
is greater than km, where m is the mean conversion tract
length and k is a positive integer, becomes negligible as k
 grows. So, only a finite sequence (say 50m) on the left of the
sequence of interest needs to be taken into account for the
latter kind of events and the rest of the chromosome can be
ignored.

Let ρ denote the population crossing-over rate, γ the pop-
ulation gene-conversion rate, and θ the population mutation
parameter. Let X denote a sample of haplotypes or genotypes
and GS denote a collection of correlated trees (i.e., an ARG)
at the marker locations consistent with X. We are interested
in sampling from the following posterior probability density:

\[ f(\rho, \gamma, m|X) = \frac{\int f(X|GS, \theta)f(GS|\rho, \gamma, m)f(\rho)f(\gamma)f(m)d\theta dGS}{f(X)}. \tag{1} \]

The posterior distribution of recombination parameters
can be numerically evaluated by using a reversible-jump
MCMC (RJMCMC) scheme. In particular, a Metropolis–Hast-
ings (MH) algorithm is used to estimate the parameters de-
scribed in Equation 1. The MH algorithm has two steps: (i)
the proposal step in which potential new parameter values
are simulated from the proposal density Q(ϕ’|ϕ) and (ii)
the acceptance step in which the proposed values are ac-
cepted with probability α or rejected with probability 1 −
α. If accepted, ϕ’ becomes the current state in the chain;
otherwise the chain remains at ϕ and ϕ’ is discarded. The
acceptance probability for a parameter is given by

\[ \alpha = \min \left( 1, \frac{l(\phi'|X)Q(\phi|\phi')f(\phi')}{l(\phi|X)Q(\phi'|\phi)f(\phi)} \right), \tag{2} \]

where f denotes the prior density and l denotes the likeli-
hood. For each node in the ARG, we store an array of size
equal to the number of markers and keep track of marker
positions that are ancestral to the sample. We call this the
ancestry vector [also the Marker Ancestry (MA) vector, see
Wang and Rannala 2008]. The distance between the start
and the end of the ancestral locations in this vector is re-
ferred to as the length of the ancestry vector. Our joint es-
 timation program begins with a binary tree that is consistent
with the observed data and then proceeds according to the
following proposal moves.

**Adding (or removing) a pair of crossing-over and
coalescence nodes**

We generate two uniform random variables u and v from the
interval (0, ti0), where t10 denotes the height of the geneal-
ogy. The smaller of the two becomes the time for inserting
a new crossing-over node and the larger is the time corre-
sponding to the coalescence node. The remainder of this
move is identical to the one in Wang and Rannala (2008)
except that the probability of acceptance becomes

\[ \alpha = \min \left( 1, \frac{l(X|GS', \theta)Q(GS|GS')f(GS'|\rho, \gamma, m)}{l(X|GS, \theta)Q(GS|GS)f(GS|\rho, \gamma, m)} \right), \]

where the proposal ratio is

\[ \frac{Q(GS|GS')}{Q(GS'|GS)} = \frac{\frac{\rho^2 n'_{br1} n'_{br2}}{2n_{pair}}(l_2 - l_1)}{\text{GS denotes the current genealogy and GS' denotes the newly
proposed genealogy.} n_{pair} \text{ here refers to the number of pos-
sible pairs of crossing-over and coalescence nodes that can
be deleted from GS'.} n_{br1} \text{ and } n_{br2} \text{ refer to the number of
eligible branches at the times at which new crossing-over
and coalescence nodes will be inserted where eligible
branches must have ancestry vector length >1 when insert-
ing crossing-over nodes. } l_2 - l_1 \text{ refers to the distance be-
tween the start and the end of the ancestral segment in a
branch.}

The probability of adding a pair of nodes to move from GS
to GS' is given by

\[ (2/t_{i0}^2) \times (1/n_{br1}) \times (1/n_{br2}) \times (1/(l_2 - l_1)). \]

The first term denotes the probability of proposing the ages
of the two nodes, the second and third terms denote the
probability that the two nodes are inserted into the
particular eligible branches at their respective times, and
the last term denotes the probability of proposing a re-
combination breakpoint in the crossing-over node that is
inserted. Note that when deleting a pair of crossing-over and
coalescence nodes, we randomly choose a pair to delete.
Thus, in the proposal ratio 1/n_{pair} denotes the probability
that the same pair of crossing-over and coalescent nodes
that were added are chosen for deletion, to move back from
GS' to GS.

**Calculating the likelihood of data given a genealogy GS:**

l(X|GS, θ) denotes the likelihood of observing a set of
haplotypes/genotypes given an ARG and the population
mutation rate. Given the genealogical tree \( \tau_i \) for a marker site \( i \), and conditional on one or more mutations having occurred, the likelihood of a marker tree is given by

\[
I(D_i \mid G_i) = \frac{(1 - e^{-0.5\rho t_i}) \pi_D + e^{-0.5\rho t_i}}{1 - e^{-0.5\rho t_i}}
\]

where \( l \) indicates a branch in \( \tau_i \) with length \( t_i \), connecting \( D_i \) and \( D_{i-k} \), with \( D_{i-k} \) being the ancestral allele of \( D_i \) in that branch. The likelihood is calculated across all branches in \( \tau_i \) and \( l \) denotes the indicator function. The parameter \( T_i \) represents the sum of the branch lengths in \( \tau_i \) and \( \pi_i \) is the frequency of allele \( j \) at stationarity. \( I(X \mid G_i, \theta) \) is the product of the likelihoods across all the marker trees.

**Adding (or removing) a pair of gene-conversion and coalescence nodes**

As before, we generate two uniform random variables \( u \) and \( v \) from the interval \((0, t_{i0})\), where \( t_{i0} \) denotes the height of the genealogy. The smaller of the two becomes the time for inserting a new gene-conversion node and the larger is the time corresponding to the coalescence node. Note that we are interested only in gene-conversion events that can potentially change the genealogy of the sample at the marker locations and thus the sample configuration.

Let \( l_2 - l_1 \) denote the distance between the start and the end of the ancestral segment in a branch of the graph. We propose a gap position \((g)\) for initiation of gene conversion nonuniformly from a region of length \( l_2 - l_1 + 50m \), where the sequence of length 50m lies to the immediate left of the start of the ancestral region in the MA vector and \( m \) denotes the mean conversion tract length.

The probability that a gap location is proposed is proportional to the chance that the tract length is greater than or equal to the distance \((k)\) from the nearest MA vector position on the right of that gap location. The chance that the tract length is \( \geq k \) is given by

\[
q(1 - q)^{k-1} + q(1 - q)^k + q(1 - q)^{k+1} + \cdots = \frac{q(1 - q)^{k-1}}{1 - (1 - q)} = (1 - q)^{k-1}.
\]

After choosing the initiation gap, we propose a tract length \((z)\) according to a truncated geometric distribution conditional on the tract length greater than or equal to the distance from the nearest MA vector position on the right, so that only tracts that can affect the genealogy of the sample at markers are included. When the start of the gene-conversion tract is to the left of the start of the MA vector, we include the additional condition that the end of the tract is always to the left of the end of the MA vector. Thus, we exclude tracts \( \geq l_2 - l_1 + k \) for such cases because such tracts do not affect the sample configuration (the probability of picking such a gap is proportional to \( (1 - q)^{k-1} - (1 - q)^{l_2 - l_1 + k - 1} \). The probability of acceptance is equal to

\[
\alpha = \min \left( 1, \frac{l(X \mid G_i, \theta)Q(G_i \mid G_i, G_i)\sum_{\gamma = 0}^{\gamma = m}(G_i \mid \rho, \gamma, m)}{l(X \mid G_i, \theta)Q(G_i \mid G_i, G_i)\sum_{\gamma = 0}^{\gamma = m}(G_i \mid \rho, \gamma, m)} \right),
\]

where the proposal ratio is

\[
\frac{Q(G_i \mid G_i, G_i)}{Q(G_i \mid G_i, G_i)} = \frac{n_{10}n_{10}n_{10}n_{10}}{2n_{10}p_{10}d(z)}
\]

The terms are the same as before except that \( p(g) \) denotes the chance that a gap location is proposed. Let \( p_g \) denote the probability that a gene-conversion tract that initiates at the base pair along the sequence includes some but not all ancestry vector positions. Then, \( p(g) = P_g / \sum_{j=1}^{j=10} p_j \). \( d(z) \) represents the density of the truncated geometric distribution for tract length at the chosen gap location; i.e., \( P[Z = z \mid \text{Tract}] \) includes some but not all MA vector positions. Note that the proposal distributions for both initiation points and tract lengths are identical to the prior distributions when gene-conversion rates are assumed to be uniform along the sequence. \( n_{10} \) refers to the number of possible pairs of gene-conversion and coalescence nodes that can be deleted. \( n_{10} \) and \( n_{10} \) refer to the number of eligible branches at the times at which new gene-conversion and coalescence nodes will be inserted, where eligible branches must have ancestry vector length \( > 1 \) when inserting gene-conversion nodes.

**Local topology rearrangements**

This move consists of moving either a recombination or a coalescence node to a new location in the graph. It is identical to what is described in Wang and Rannala (2008) except that the conditional probability terms will include parameters \( \rho \), \( \gamma \), and \( m \) instead of \( \rho \) alone.

**Propose new waiting times between events in the graph**

We propose a new graph with new waiting times between the successive (recombination or coalescence) events. The topology of the graph is left unchanged in this new move. The waiting times to the next event are proposed from the prior distribution expected under coalescent theory on the basis of the number of lineages and the ancestry vectors after the previous event. Note that this proposal is independent of the current waiting times. The probability of acceptance is given by

\[
\alpha = \min \left( 1, \frac{l(X \mid G_i, \theta)Q(G_i \mid G_i, G_i)\sum_{\gamma = 0}^{\gamma = m}(G_i \mid \rho, \gamma, m)}{l(X \mid G_i, \theta)Q(G_i \mid G_i, G_i)\sum_{\gamma = 0}^{\gamma = m}(G_i \mid \rho, \gamma, m)} \right),
\]

Let \( k \) denote the number of lineages after the previous event. Under coalescent theory, the rate of coalescence is \( 0.5k(k - 1) \), the rate of crossing over is \( 0.5kp \), the rate of gene conversion is \( 0.5ky \), and the waiting time to the next event is exponentially distributed with an overall rate of \( [0.5k(k - 1) + 0.5kp + 0.5ky] \). However, because here we are interested only in informative gene-conversion and
crossing-over events, we have to make some modifications when calculating the rates. Let \( f_i \) denote the fraction of the sequence included within the start and the end of the ancestry vector. Then, the rate of informative crossing-over events for the \( i \)th lineage is given by \( p_i \) and the overall rate of crossing over becomes \( p_t = \sum_{i=1}^{n} p_i \). Similarly, let \( p_i \) denote the probability that a gene-conversion tract that initiates at the \( j \)th basepair along the sequence in lineage \( i \) includes some but not all ancestry vector positions at that lineage. Let \( \gamma / L \) denote the gene-conversion rate per base pair. Then, the rate of informative gene conversions is given by \( \gamma_t = \sum_{i=1}^{n} \sum_{j=1}^{n} l_1(i) - l_2(i) - 50m p_i \gamma / L \), where \( l_1(i) \) and \( l_2(i) \) denote the start and the end of the ancestry vector for the \( i \)th lineage. Thus, the waiting times to informative events are exponentially distributed with a rate of \( [0.5(k - 1) + 0.5p_t + 0.5\gamma_t] \). The overall probability of a proposal is equal to the product of the exponential density functions of successive events. Note that since we propose the new waiting times from the coalescent prior,
\[
\frac{Q(G_i|G_j) f(G_i'|p, \gamma, m)}{Q(G_i'|G_j) f(G_i|p, \gamma, m)} = 1.
\]

**Coalescent prior density for a genealogy:** To calculate the coalescent prior probability for a genealogy \( G_S \), i.e., \( f(G_S|p, \gamma, m) \), we calculate the product of three quantities at each event and multiply them together:

1. Density of the waiting time for that event: The time is exponentially distributed with a rate of \( [0.5(k - 1) + 0.5p_t + 0.5\gamma_t] \).
2. The probability that the event is a coalescence or a gene conversion or a crossing over:
\[
\begin{align*}
P(\text{Coalescence}) &= \frac{0.5k(k - 1)}{[0.5(k - 1) + 0.5p_t + 0.5\gamma_t]} \\
P(\text{gene conversion}) &= \frac{0.5\gamma_t}{[0.5(k - 1) + 0.5p_t + 0.5\gamma_t]} \\
P(\text{crossing over}) &= \frac{0.5p_t}{[0.5(k - 1) + 0.5p_t + 0.5\gamma_t]}
\end{align*}
\]
3. The chance is calculated that (a) a particular pair of lineages will be picked to coalesce; i.e., \( 2/k(k - 1) \) if the event is coalescence. (b) If the event is crossing over, the chance that a particular lineage will be picked for crossing over is multiplied by the chance that its breakpoint will be picked. (c) If the event is gene conversion, the chance that the particular lineage is picked for gene conversion is multiplied by the chance that its initiation point is chosen multiplied by the chance that the tract length is chosen.

**Propose a new breakpoint (and tract length) for a recombination node**

In this move, we pick a crossing-over or gene-conversion node at random and propose a new breakpoint (and tract length) for that node. We also randomly propose a new direction for how the products of recombination get distributed (i.e., which part goes to the left branch/right branch of the recombination node). We then update the ancestry vectors of the subsequent nodes in the graph on the basis of the modified node. The probability of acceptance as before is equal to
\[
\alpha = \min \left( 1, \frac{l(X|G_{S^*}, \theta) Q(G_{S^*}|G_S) f(G_{S^*}|p, \gamma, m)}{l(X|G_S, \theta) Q(G_S|G_{S^*}) f(G_S|p, \gamma, m)} \right)
\]

If the chosen node is a crossing-over node, we propose a new break point uniformly between the start and the end of the ancestry vector positions and the proposal ratio for the node is equal to 1. The proposal ratio for a new breakpoint and tract length for a gene-conversion node is given by
\[
\frac{Q(G_S|G_{S^*})}{Q(G_{S^*}|G_S)} = \frac{p(g)d(z)}{p(g')d(z')}
\]

Here, \( p(g) \) denotes the probability that a breakpoint location is proposed in a gene-conversion node. Let \( p_i \) denote the probability that a gene-conversion tract that initiates at the \( j \)th basepair along the sequence includes some but not all ancestry vector positions. Then, \( p(g) = p_i \sum_{i=1}^{n} l_1(i) - 50m p_i \gamma / L \), where \( l_1(i) \) and \( l_2(i) \) denote the start and the end of the ancestry vector, respectively. \( d(z) \) is the density of the truncated geometric distribution for tract length at the chosen gap location; i.e., \( P(Z = z) \) Tract includes some but not all MA vector positions.

**Modify ancestral states, haplotype phases, and missing data**

These steps are identical to those found in Wang and Ranala (2008) because the likelihood of the data given the genealogy trees and \( \theta \), i.e., \( l(X|G_s, \theta) \), is the same as described there.

**Propose new values for the parameters \( p, \gamma, m, \) and \( \theta \)**

New \( p \) is proposed according to a sliding window with reflecting boundary at 0. The acceptance probability is given by
\[
\alpha = \min \left( 1, \frac{f(G_S|p', \gamma, m) f(p')}{f(G_S|p, \gamma, m) f(p)} \right)
\]

New \( \gamma \) is proposed according to a sliding window with reflecting boundary at 0. The acceptance probability is given by
\[
\alpha = \min \left( 1, \frac{f(G_S|p, \gamma', m) f(\gamma')}{f(G_S|p, \gamma, m) f(\gamma)} \right)
\]

New \( m \) is proposed according to a sliding window with reflecting boundaries at 1 and at 2000 or 3000 bp. The acceptance probability is given by
\[ \alpha = \min \left( 1, \frac{f(G_S|\rho, \gamma, m') f(m')}{f(G_S|\rho, \gamma, m) f(m)} \right). \]

New $\theta$ is proposed according to a sliding window with reflecting boundary at 0. The acceptance probability is given by
\[ \alpha = \min \left( 1, \frac{l(X|G_S, \theta') f(\theta')}{l(X|G_S, \theta) f(\theta)} \right). \]

**Variable recombination rate model**

We also use a variable recombination rate model in our analysis that includes background rate variation and hotspots. In this model, the background crossing-over rate follows a gamma distribution with shape and scale parameters and recombination hotspots that arise according to a Markov process. The waiting distance until the occurrence or loss of a crossing-over hotspot is exponentially distributed while the intensity of a hotspot follows a log-normal distribution. More details about this crossing-over model can be found in Wang and Rannala (2009). In our extension of the Wang and Rannala model, we assume that gene-conversion and crossing-over rates vary in an identical pattern such that their ratio ($f$) remains uniform along the sequence. Thus, all crossing-over hotspots are also gene-conversion hotspots.

**Results**

**Checking the MCMC program**

To check the correctness of our MCMC algorithm, we first examined the stationary distribution of several genealogy-based summary statistics when the chain is run with no data (i.e., with the likelihood ratio set to 1). We fixed the values of $\rho$, $\gamma$, and $m$ when running the chain and then compared the distribution of the number of crossing-over (CO) events, the number of gene-conversion (GC) events, and the genealogy height ($H$) with coalescent simulations for the same parameters. As can be seen from Table 1, the expectations from the MCMC algorithm are in agreement with those obtained by straightforward Monte Carlo simulations under the coalescent.

**Test runs on simulated data:** Next, we tested our inference program on data sets simulated with gene conversion alone and compared the posterior distribution of $\gamma$ as estimated by our program with the true value used in simulations. For this testing, we fixed the mean tract length and $\theta$ to their true values and $\rho$ was fixed at 0. The results obtained are shown in supporting information, Figure S1. When gene-conversion rates are high, we expect that the power to detect gene conversion will be high and our posterior distribution will not include 0 most of the time (e.g., $\gamma = 4000/\text{Mb}$, Figure S1), which would tell us clearly that conversions have happened. On the other hand, when gene-conversion rates become lower it becomes difficult to infer whether a conversion event has happened in a particular sample’s history (because the posterior will often include 0). This is because not every gene-conversion event is necessarily detectable in a sample and with only a few conversion events in a sample history, by chance, we could have samples that are also consistent with one or more binary trees.

In Figure S1ii we show results when we run our joint estimation program on data sets simulated with gene conversion alone with the tract length fixed at its true value. We can see that estimates of $\rho$ are low but nonzero and the credible intervals contain values close to 0. Note that estimates of $\gamma$ under this joint estimation scenario can be considerably lower than the truth although the true value of $\rho$ is 0 (which means that we interpret some of the gene conversions as crossing overs). In Figure S1iii we show results for the other boundary case, i.e., when data are simulated with crossing over alone with $\gamma$ set at 0. For this scenario, we attempted to estimate both crossing-over and gene-conversion rates with the tract length fixed at different values. In addition, we also estimated the recombination parameters and mean tract lengths jointly. We can see that estimates of $\gamma$ can be substantial although the true value is 0. When the mean conversion tract length becomes longer (e.g., 2000 bp, 8000 bp), we observe multiple peaks in the joint posterior distribution. This suggests that more than one combination of crossing-over and gene-conversion rates can explain the simulated data and there is difficulty in distinguishing between the effects of these two processes.

For Figure S2, we assumed that both gene-conversion rates and mean tract lengths are unknown and estimated them jointly. Subsequently, we performed simulations with both crossing over and gene conversion and attempted to jointly estimate both $\gamma$ and $\rho$ for these data sets with the mean tract length set to its true value. The corresponding results are shown in Figure S3. In all the figures (i.e., Figure S1, Figure S2, Figure S3), we can see that the modes of the posterior distribution are close to the true values of the recombination parameters. In Figure S3 too, we sometimes see more than one peak in the posterior distribution.

In general, estimating gene-conversion rates from population genomic data is a challenging problem. This is because the traces of this phenomenon in data are limited and subtle. Note that the ability to detect and quantify gene conversion in both experiments and population genetic analysis depends on the tract size. When tracts are short compared to the spacing between markers, a substantial fraction of gene-conversion events will not leave a trace in the sample, making it difficult to infer rates accurately. On the other hand, when tracts are long compared to the sequence of interest, some fraction of them will fall only partly inside the sequence and will be indistinguishable from crossing over. As Figure S1, ii and iii, and Figure S3 highlight, there can be substantial confounding when attempting to jointly estimate both these parameters and it is difficult to tease apart the effects of conversion and crossing over from one another.
Simulations with variable recombination rates: We tested an implementation of InferRho that incorporates variation in crossing-over and gene-conversion rates on data sets simulated with recombination hotspots. We simulated 20 samples of 20-kb sequences with $\theta = 20.0$, $m = 500$ bp, and a single hotspot between 9 kb and 11 kb. Both $\rho$ and $\gamma = 50$/Mb outside hotspots, whereas within the hotspot these numbers were 5000/Mb. The ratio of gene conversion to crossing over ($f$) was assumed to be uniform along the sequence in these simulations with $f = 1.0$. Table 2 shows the results obtained from running our program on three independent replicates for these parameters.

Next, we simulated 50 samples of 20-kb sequences with $\theta = 20.0$, $m = 500$ bp, a single hotspot between 9 kb and 11 kb, and uniform $f = 10.0$ (parameters used in Gay et al. 2007). $\rho = 50$/Mb and $\gamma = 500$/Mb outside hotspots, whereas within the hotspot these numbers were 5000/Mb and 50,000/Mb, respectively. We estimated recombination rates for these data sets using the program of Gay et al. (2007), which implements a variable gene-conversion rate model. Note that this program does not try to identify hotspots but estimates a different crossing-over and gene-conversion rate between adjacent SNPs. We also inferred the locations of hotspots using InferRho. Figure 1 plots the results obtained using these two different methods on five independent replicates for these parameters. In both Table 2 and Figure 1, we can see that the estimated locations of the starts and ends of hotspots are close to the true locations.

Comparison with other methods: We compared the performance of our method (implemented in the software package InferRho) with that of the methods described in Gay et al. (2007) and Yin et al. (2009). For this comparison, we simulated 100 data sets each for two different parameter combinations and estimated all three recombination parameters (i.e., the population crossing-over rate, the population gene-conversion rate, and the mean tract length) jointly for these data sets. Then, we calculated summaries of these estimates and compared them with those obtained from the other two methods. These comparisons are shown in Table 3. Note that the summaries for the methods of Gay et al. (2007) and Yin et al. (2009) are based on the same set of simulated data sets whereas those for our method are based on an independent sample of 100 data sets. When implementing all three methods, the starting values of the recombination parameters were set to the true value that was used in simulations.

For all the parameters tested, we find that the performance of the InferRho program and the method of Yin et al. (2009) are better than that of Gay et al. (2007). Presumably this is because the approach of Gay et al. (2007) ignores overlapping gene-conversion events while the other two methods do not assume this restriction. In theory, we would expect the Gay et al. (2007) method to perform worse if overlapping gene-conversion events are significantly frequent in the history of the sample. Compared to the method of Yin et al. (2009), InferRho appears to have higher mean squared error for tract length estimation but lower mean squared error for crossing-over and gene-conversion rate estimation. This may be due to confounding when attempting to estimate both gene-conversion rates and tract lengths jointly from population genetic data.

Estimates in real data

We applied our method to jointly estimate the crossing-over and gene-conversion rates in two genes, the $su-w$ and $su-s$ genes of D. melanogaster that were first studied by Langley et al. (2000). These genes are located close to the telomeric

Table 1 Comparison between the MCMC program and coalescent simulations

| $\rho^a$ | $\gamma^a$ | $E(\text{CO})^b$ | $E(\text{CO})^c$ | $E(\text{GC})^b$ | $E(\text{GC})^c$ | $E(H)^b$ | $E(H)^c$ | Mean tract | Sample size | $M^d$ | $L^e$ |
|---------|-----------|------------------|------------------|------------------|------------------|----------|----------|------------|-------------|-------|-------|
| 0.000   | 0.300     | 0.0000           | 0.0000           | 4.4084           | 4.4233           | 2.4454   | 2.4572   | 5          | 8           | 4     | 4     |
| 0.000   | 0.100     | 0.0000           | 0.0000           | 1.2583           | 1.2614           | 1.9721   | 1.9781   | 2          | 8           | 4     | 4     |
| 0.100   | 0.100     | 0.3163           | 0.3162           | 0.5782           | 0.5777           | 1.2694   | 1.2682   | 5          | 2           | 4     | 4     |
| 0.300   | 0.300     | 2.4264           | 2.4281           | 2.7716           | 2.7760           | 4.4577   | 4.4683   | 5          | 8           | 4     | 4     |
| 0.100   | 0.300     | 0.8060           | 0.8061           | 2.5602           | 2.5605           | 4.4287   | 4.4282   | 5          | 8           | 4     | 4     |
| 0.001   | 0.001     | 29.884           | 30.646           | 3.3402           | 3.4341           | 5.7887   | 5.9864   | 100        | 8           | 8     | 8000  |

Notes: 

$^a$ Population crossing-over and gene-conversion rates per base pair.

$^b$ Average of crossing-over count, gene-conversion count, and height of genealogy from coalescent simulations.

$^c$ Average of crossing-over count, gene-conversion count, and height of genealogy from MCMC algorithm.

$^d$ Number of markers.

$^e$ Sequence length in base pairs.

Table 2 Estimated locations of hotspots in three independent simulated data sets

| Data set | Hotspot start, bp | Hotspot end, bp | Hotspot intensity estimate | $f$ |
|----------|-------------------|-----------------|---------------------------|-----|
| Replicate 1 | 9,486 (8,746–10,226) | 10,226 (9,240–11,557) | 4,030.71 | 2.375 (0.375–6.125) |
| Replicate 2 | 10,885 (5,240–16,530) | 11,557 (9,240–11,068) | 4,030.71 | 0.375 (0.125–7.375) |
| Replicate 3 | 9,822 (8,087–11,557) | 11,068 (9,240–11,879) | 3,968.12 | 3.625 (1.125–4.625) |

Each simulated data set consists of 20-kb sequences and 20 samples with a single hotspot of intensity 5000/Mb between 9 kb and 11 kb. The relative rate of gene conversion to crossing over ($f$) was uniform along the sequence and was equal to 1. The estimated locations for the start and end of hotspots and $f$ are shown along with the 90% credible intervals in parentheses.
regions of the X chromosome and are \(~4.1\) kb and \(2.5\) kb long, respectively. The \(su-s\) data set contains 50 haplotypes and 41 SNPs whereas the \(su-w\) data set that we used contains 50 haplotypes and 46 SNPs in an African-only sample. We did not try to estimate tract lengths in our initial analysis and fixed them to 352 bp (estimates from Hilliker et al. 1994). The value of \(\theta\) was set to be equal to Watterson’s estimate for each data set. Table 4 shows the results obtained for this analysis (see also Figure S5).

Subsequently, we implemented a version of our inference program where we estimated both mean tract lengths and recombination rates jointly. These results are shown in Table 5. Figure S4 shows 95% credible intervals with maximum posterior density for the estimated parameter values. Note that the estimated rate of gene conversion depends on the mean tract length and for shorter tracts the inferred rates are expected to be higher (see also Figure S5).

We also compared our results with those obtained by the methods in Gay et al. (2007) and Yin et al. (2009) in Tables 4 and 5. We observe that there is some difference between the point estimates provided by the three different programs. Presumably, this is because there is limited
information in these data sets for joint estimation and there is difficulty in distinguishing between the effects of the different recombination mechanisms in the short sequences of the Drosophila genes.

Bayes factor in real data

We compared the posterior distribution of gene-conversion start points in the su-w and su-s genes with the distribution under an uninformative prior. The prior distributions were obtained by running the chain without any data (i.e., with likelihood ratio = 1) for the same sample size and marker positions as in the real data sets and assuming uniform recombination rates. Let $p_1$ denote the probability that a conversion initiation point lies within a particular window for the prior sample of genealogies (This is the fraction of initiation points that lie within that window). Similarly, let $p_2$ denote the corresponding probability for the posterior sample of genealogies. The odds for the prior and posterior for a window are given by $p_1/(1 - p_1)$ and $p_2/(1 - p_2)$, respectively. Formally, the Bayes factor (BF) is defined as the ratio of the posterior and prior odds; i.e.,

$$BF = \frac{p_2(1 - p_1)}{p_1(1 - p_2)}.$$

The log of the BF should be interpreted as the change in the evidence for gene-conversion initiation in a region due to the data. A large negative value indicates evidence against gene-conversion initiation and a large positive value indicates evidence for gene-conversion initiation. The BF for 10-bp windows is plotted along the genes, assuming the mean tract length = 352 bp (see Figure 2, A and B). Note that the Bayes factors for the windows were expected to indicate where gene-conversion events have occurred in the history of the sample but their values do not necessarily imply high or low rates. Thus, this analysis is unrelated to hotspot identification described in the Simulations with variable recombination rates section. For these Drosophila data sets, it seems that there is little information concerning the location of gene-conversion start points because we do not see very large or very small BF s ($|\log(BF)| < 2$ generally).

Discussion

We have described a Bayesian MCMC method for jointly inferring the crossing-over and gene-conversion parameters from SNP data sets that extends the crossing-over estimation method originally proposed by Wang and Rannala (2008). In this method, we model the genealogy of a sample as a recombination graph and keep track of ancestral marker sites in the MA vectors of its nodes. Furthermore, we include only informative crossing-over and gene-conversion events that can potentially change the genealogy of the sample at the marker locations. These aspects of our algorithm make the full-likelihood estimation of recombination rates efficient in terms of both the running time and memory requirements. Assuming an uninformative prior for the parameters of interest, we can estimate their joint posterior distribution using a reversible-jump MCMC scheme. In the Metropolis–Hastings step, we propose changes to various features of the

Table 3 Comparison between three different methods for recombination rate estimation

| Method       | $\rho^a$ | $\gamma^a$ | $m^a$ | MSE($\rho$)$^b$ | MSE($\gamma$)$^b$ | MSE($m$)$^b$ | #($\rho$):2)$^c$ | #($\gamma$):2)$^c$ | #($m$):2)$^c$ |
|--------------|----------|------------|-------|-----------------|-----------------|---------------|-----------------|-----------------|-----------------|
| Gay et al. (2007) | 500   | 500         | 500   | 3.488           | 1,022,389       | 0.723         | 0.71            | 0.26            | 0.36            |
| Yin et al. (2009) | 500   | 500         | 500   | 0.079           | 0.661           | 0.105         | 0.71            | 0.55            | 0.88            |
| InferRho      | 500   | 500         | 500   | 0.041           | 0.222           | 0.644         | 0.89            | 0.77            | 0.56            |
| Gay et al. (2007) | 500   | 1000        | 500   | 0.085           | 618,219         | 0.349         | 0.78            | 0.48            | 0.35            |
| Yin et al. (2009) | 500   | 1000        | 500   | 0.060           | 1,348           | 0.098         | 0.79            | 0.70            | 0.87            |
| InferRho      | 500   | 1000        | 500   | 0.043           | 0.487           | 0.250         | 0.88            | 0.81            | 0.62            |

$^a$ $\rho$ denotes population crossing-over rate, $\gamma$ denotes the population gene-conversion rate, and $m$ denotes the mean conversion tract length for the 100 simulated data sets. Recombination rates and mean tract lengths are shown in units of per megabase and base pair, respectively.

$^b$ MSE, the mean square error of the parameter estimates for the 100 simulated data sets $\times 10^{-6}$. For InferRho, we calculated the marginal estimate of each recombination parameter from the joint posterior distribution. Numbers for the other methods are taken from Yin (2010).

$^c$ #($\rho$):2) represents the fraction of the 100 data sets for which estimates are within a factor of 2 of the true parameter value used in simulations (i.e., $\rho$). #($\gamma$):2) and #($m$):2) are defined similarly.
ancestral recombination graph such as the relative locations of the recombination or coalescence nodes, the number of crossing-over or gene-conversion nodes, the ancestral alleles, and the waiting times between consecutive events in the graph, etc. The method has been implemented in a new version of the software package InferRho (see Wang and Rannala 2008), which will become available online at http://rannala.org. Currently, we are checking the software’s performance for larger values of \( \rho \) and \( \gamma \) (\( \approx 50.0 \)).

The MCMC method described here attempts to generate a posterior sample of genealogies for the data under the coalescent. It uses all the information contained in the haplotypes or genotypes for this purpose (because we propose and accept only genealogies consistent with the observed data) and theoretically, after a very large number of iterations, we expect samples from essentially the correct posterior distribution. In contrast, PAC-based models (e.g., Hellenthal 2006; Gay et al. 2007; Yin et al. 2009) evaluate likelihoods of the data assuming a heuristic model that is an approximation of the constant population size coalescent. The relationship between the PAC model and the coalescent process is not well understood theoretically although they appear to be analogous. Apart from the benefit of having a posterior sample of genealogies, the main advantage of our Bayesian approach compared to other alternatives is that we expect to obtain better confidence intervals for the estimated parameter values (if the coalescent is a closer approximation to reality). It is also worth mentioning that the PAC model developed by Yin et al. (2009) considers only a restricted class of gene-conversion events (with regard to overlaps between events) whereas our method is more general and models all the conversion events that can potentially affect the configuration of the sample. Finally, since the InferRho algorithm lies within the coalescent framework, it is flexible and in principle can be extended for estimation under demographic models other than constant sized ones (e.g., exponential growth, population bottlenecks, etc.).

We performed two different kinds of tests on simulated data to verify the correctness of our estimation program. First, we ran the program without any data using a likelihood ratio of 1 and compared the average of several genealogy-based summary statistics to those obtained by Monte Carlo simulations under the coalescent. Second, we simulated data sets for which the true parameter values are known and compared them with the mode of the posterior distributions as estimated by the program. From both these tests, it appears that the modified InferRho program is working correctly. In addition to these, we also compared the accuracy of our inference method with that of two other comparable methods (proposed in Gay et al. 2007 and Yin et al. 2009) for samples of 100 simulated data sets. For estimating crossing-over and gene-conversion rates, InferRho has smaller mean squared error compared to the methods of Gay et al. (2007) and Yin et al. (2009) for the parameters tested in this study. For estimation of tract length, the mean squared error of InferRho is lower than the program of Gay et al. (2007) but higher than that of Yin et al. (2009).

We used the new version of InferRho to jointly estimate crossing-over and gene-conversion rates in the \( su-w \) and \( su-s \) genes of \( D. \) melanogaster. Our analysis with a fixed mean tract length of 352 bp suggests that gene conversion occurs more frequently than crossing over in these regions while the local estimates of crossing-over rate are not low. Furthermore, we also attempted to jointly estimate recombination rates and gene-conversion tract lengths and from this analysis it appears that mean tract lengths are \(~70\) bp and \( 430\) bp for the \( su-w \) and \( su-s \) genes, respectively. Finally, we compared the posterior distribution of gene-conversion initiation points with an uninformative prior distribution, using the Bayes factor calculated for \( 10\)-bp windows along these genes. Plots of the logarithm of Bayes factors for gene-conversion start points do not indicate any strong deviations from a prior distribution expected under the uniform recombination rate model. This suggests that there is not sufficient information in the data to infer the locations of historical conversion breakpoints.

### Table 4 Joint estimates of crossing over (\( \rho \)) and gene conversion (\( \gamma \)) for \( m = 352\) bp

| Gene | Method       | \( \rho/\)Mb per Mb | \( \gamma/\)Mb per Mb | \( \hat{t} = \gamma/\rho \) |
|------|--------------|---------------------|-----------------------|------------------------|
| su-s | Gay et al. (2007) | 1,700               | 12,000                | 7.10                   |
| su-s | Yin et al. (2009) | 2,240               | 11,510                | 5.14                   |
| su-s | InferRho     | 1,500 (700–5,500)   | 6,900 (3,500–10,300)  | 4.60 (0.875–7.875)     |
| su-w | Gay et al. (2007) | 570                 | 28,000                | 48.0                   |
| su-w | Yin et al. (2009) | 33                  | 27,040                | 819.4                  |
| su-w | InferRho     | 4,300 (2,500–8,100) | 11,700 (5,300–16,100) | 2.72 (0.875–3.625)     |

The 95% maximum posterior density credible intervals for InferRho are shown in parentheses. For InferRho, MCMC chains were run for 40 million iterations and the first 10 million iterations were discarded as burn-in. For other methods, the numbers are taken from Yin (2010).

### Table 5 Estimates of crossing over (\( \rho \)), gene conversion (\( \gamma \)), and mean tract length (\( \mu \))

| Gene  | Method       | \( \rho/\)Mb | \( \gamma/\)Mb | \( m, \) bp |
|-------|--------------|-------------|-------------|-----------|
| su-s  | Gay et al. (2007) | 920         | 11,600      | 480       |
| su-s  | Yin et al. (2009) | 1,290       | 9,860       | 560       |
| su-s  | InferRho     | 3,350       | 5,950       | 430       |
| su-w  | Gay et al. (2007) | 8,520       | 251,130     | 5         |
| su-w  | Yin et al. (2009) | 1,450       | 41,090      | 162       |
| su-w  | InferRho     | 5,550       | 28,950      | 70        |

For InferRho, MCMC chains were run for 41 million and 37 million iterations (for \( su-s \) and \( su-w \), respectively) and the first 10 million iterations were discarded as burn-in. We show the marginal estimate of each parameter from the joint posterior distribution. For other methods, the numbers are taken from Yin (2010).
One area of concern with using full-likelihood methods is their high computational expense when data sets become large. Note that the execution time of the InferRho algorithm is much higher (orders of magnitude) than those of Gay et al. (2007) and Yin et al. (2009). Therefore, it would be desirable to make such inference methods as optimal as possible in terms of run-time efficiency. To improve speed, we implemented several optimizations in the current version of our program. In our MA vectors, we keep track of marker locations that are ancestral to the sample and exclude uninformative gene-conversion and crossing-over events in the ARGs. We use arrays of size equal to the number of markers to represent the MA vectors. Operations on MA vectors (e.g., merging two MA vector arrays during coalescence, finding which marker positions have coalesced in a node, etc.) take linear time in terms of the total number of markers. Second, we use lookup tables to keep track of the coalescent prior likelihood of a graph and the likelihood of observed data given a graph once they have been calculated. Since a majority of newly proposed genealogies do not get

Figure 2. (A) Bayes factors along the su-w gene for 10-bp windows. The logarithm of the ratio of posterior to prior odds for gene-conversion initiation points is plotted along the gene. The su-w gene is 2.5 kb long. (B) Bayes factors along the su-s gene for 10-bp windows. The logarithm of the ratio of posterior to prior odds for gene-conversion initiation points is plotted along the gene. The su-s gene is 4.1 kb long.
accepted, these values are reused multiple times when calculating the acceptance probabilities in the algorithm. Finally, modifying some part of the graph (e.g., adding, deleting, or moving a node) during a proposal entails that we update the MA vectors of the subsequent nodes in the graph. While implementing this step, we first mark all the nodes that are ancestral to the modified node and then update only the MA vectors of the marked nodes. This previous step as well as parallelization of the code when running multiple MCMC chains can lead to further improvements in run-time efficiency.

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Bayesian Population Genomic Inference of Crossing Over and Gene Conversion

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Supporting Information
http://www.genetics.org/content/suppl/2011/08/12/genetics.111.130195.DC1
Figure S1  Estimates of gene-conversion in data simulated with conversion alone. Simulations of 10 kb sequences with $\theta = 10.0$ and mean tract ($m$) = 100, 352 or 500 bp. Each figure shown represents the posterior distribution of the population gene-conversion rate ($\gamma$) for an independent dataset with the population crossing-over rate ($\rho$) set to 0 and the tract length ($m$) set to its true value.

a) $\gamma = 4000$, $m = 500$ bp, Samples = 20  
b) $\gamma = 4000$, $m = 500$ bp, Samples = 20

c) $\gamma = 4000$, $m = 500$ bp, Samples = 20  
d) $\gamma = 2000$, $m = 100$ bp, Samples = 20
e) \( \gamma = 2000, m = 100 \text{ bp}, \text{Samples} = 20 \)

f) \( \gamma = 2000, m = 500 \text{ bp}, \text{Samples} = 20 \)

g) \( \gamma = 3000, m = 352 \text{ bp}, \text{Samples} = 20 \)

h) \( \gamma = 1000, m = 352 \text{ bp}, \text{Samples} = 20 \)

i) \( \gamma = 3000, m = 352 \text{ bp}, \text{Samples} = 20 \)

j) \( \gamma = 1000, m = 352 \text{ bp}, \text{Samples} = 20 \)
k) $\gamma = 400, m = 100 \text{ bp}, \text{Samples} = 20$

l) $\gamma = 2000, m = 500 \text{ bp}, \text{Samples} = 50$

m) $\gamma = 400, m = 500 \text{ bp}, \text{Samples} = 50$

n) $\gamma = 6000, m = 100 \text{ bp}, \text{Samples} = 40$

o) $\gamma = 6000, m = 100 \text{ bp}, \text{Samples} = 40$

p) $\gamma = 1000, m = 100 \text{ bp}, \text{Samples} = 40$
q) \( \gamma = 2000, m = 100 \) bp, Samples = 40
ii) Marginal distribution of crossing-over for 10 kb sequences simulated with gene-conversion alone and with $\theta = 10.0$. Each figure corresponds to an independent dataset and the red stars mark the 95% credible set with the maximum posterior density. The population crossing-over rate ($\rho$) and the population gene-conversion rate ($\gamma$) were jointly estimated with $m$ fixed at its true value for each dataset.

a) $\gamma = 1000$, $m = 352$ bp, Samples = 20

b) $\gamma = 3000$, $m = 352$ bp, Samples = 20
c) $\gamma = 2000$, $m = 500$ bp, Samples = 40
d) $\gamma = 6000$, $m = 100$ bp, Samples = 40
iii) Estimates of recombination parameters for 10 kb sequences simulated with crossing-over alone. Each figure corresponds to an independent dataset. The population crossing-over rate (ρ) and the population gene-conversion rate (γ) were jointly estimated for each dataset. The mean tract length (m) was either fixed to its true value or treated as a free parameter. θ was treated as a parameter and was not fixed to the true value for this analysis.

a) ρ = 1000.0, Samples = 40, θ = 10.0, m = 100 bp fixed

b) ρ = 1000.0, Samples = 40, θ = 10.0, m = 500 bp fixed
c) $\rho = 1000.0$, Samples = 40, $\theta = 10.0$, $m = 2000$ bp fixed

d) $\rho = 1000.0$, Samples = 40, $\theta = 10.0$, $m = 8000$ bp fixed
e) $\rho = 1000.0$, Samples = 40, $\theta = 10.0$
f) $\rho = 2000.0$, Samples = 20, $\theta = 10.0$
g) $\rho = 3000.0$, Samples = 40, $\theta = 10.0$
**Figure S2** Estimates of gene-conversion rate and tract length in 10 kb sequences simulated with conversion alone. Each figure represents the joint posterior distribution of population gene-conversion rate ($\gamma$) and mean tract length ($m$) for an independent simulated dataset with the population crossing-over rate ($\mu$) fixed at 0. The starting value of the parameters in the MCMC chain were set at the true values used in simulations.

a) $\gamma = 3000$, $m = 352$ bp, $\theta = 10.0$, Samples = 20
b) $\gamma = 3000, m = 352$ bp, $\theta = 10.0$, Samples = 20
c) $\gamma = 5000$, $m = 150$ bp, $\theta = 15.0$, Samples = 20
d) $\gamma = 2000$, $m = 500$ bp, $\theta = 15.0$, Samples = 20
e) $\gamma = 1000$, $m = 800$ bp, $\theta = 10.0$, Samples = 50
**Figure S3**  Estimates in data simulated with both crossing-over and gene-conversion. Each figure represents the joint posterior distribution of the population crossing-over rate ($\rho$) and the population gene-conversion rate ($\gamma$) for an independent dataset with $m$ fixed at its true value.

a) $\rho = 1000, \gamma = 1000, \theta = 10.0, m = 352$ bp, Samples = 20, 10 kb sequences.

![Graph for a)](image)

b) $\rho = 2000, \gamma = 2000, \theta = 10.0, m = 352$ bp, Samples = 20, 10 kb sequences.

![Graph for b)](image)
c) \( \rho = 3000, \gamma = 3000, \theta = 10.0, m = 352 \) bp, Samples = 20, 10 kb sequences.
d) $\rho = 1000$, $\gamma = 1000$, $\theta = 20.0$, $m = 352$ bp, Samples = 20, 10 kb sequences.
e) $\rho = 545$, $\gamma = 2700$, $\theta = 20.0$, $m = 500$ bp, Samples = 50, 20 kb sequences.

f) $\rho = 1545$, $\gamma = 2700$, $\theta = 20.0$, $m = 500$ bp, Samples = 50, 20 kb sequences*.

*\( \theta \) was not fixed at truth for this dataset but treated as a parameter when running the chain.
Figure S4  Distribution of recombination parameters and mean conversion tract length and maximum posterior density credible interval set for su-w and su-s genes. The regions included within the approximate 95% credible intervals are marked by red stars.
Figure S5  Joint posterior distribution of crossing-over and gene-conversion rates in the su-w and su-s genes of *Drosophila melanogaster* and in simulations. Each figure represents the joint posterior distribution of the population gene-conversion rate ($\gamma$) and population crossing-over rate ($\mu$) for an independent dataset with the tract length ($m$) fixed at 352 bp.

a) su-w gene dataset.

b) Simulated datasets of same length, sample size and segregating sites with rates equal to estimates for su-w gene.
c) su-s gene dataset.

d) Simulated datasets of same length, sample size and segregating sites with rates equal to estimates for su-s gene.