Gene flow biases population genetic inference of recombination rate

Kieran Samuk 1,2+, Mohamed A.F. Noor 1*

1Department of Biology, Duke University, Durham, NC 27708, USA,
2Department of Evolution, Ecology, and Organismal Biology, The University of California, Riverside, Riverside, CA 92521, USA

*Corresponding author: Department of Evolution, Ecology, and Organismal Biology, The University of California, Riverside, Riverside, CA 92521, USA. Email: ksamuk@ucr.edu

Abstract

Accurate estimates of the rate of recombination are key to understanding a host of evolutionary processes as well as the evolution of the recombination rate itself. Model-based population genetic methods that infer recombination rates from patterns of linkage disequilibrium in the genome have become a popular method to estimate rates of recombination. However, these linkage disequilibrium-based methods make a variety of simplifying assumptions about the populations of interest that are often not met in natural populations. One such assumption is the absence of gene flow from other populations. Here, we use forward-time population genetic simulations of isolation-with-migration scenarios to explore how gene flow affects the accuracy of linkage disequilibrium-based estimators of recombination rate. We find that moderate levels of gene flow can result in either the overestimation or underestimation of recombination rates by up to 20–50% depending on the timing of divergence. We also find that these biases can affect the detection of interpopulation differences in recombination rate, causing both false positives and false negatives depending on the scenario. We discuss future possibilities for mitigating these biases and recommend that investigators exercise caution and confirm that their study populations meet assumptions before deploying these methods.

Keywords: recombination; population genetics; gene flow; linkage disequilibrium; methods

Introduction

Recombination rate, the number of crossovers per unit genome per generation, plays a key role in shaping evolutionary processes and diversity in the genome. For example, through the action of linked selection, local rates of recombination are a chief determinant of patterns of genetic diversity throughout the genome (Begun and Aquadro 1992; Haddrill et al. 2014; Burri 2017; Cutter 2019; Korunès et al. 2021). Genome-wide rates of recombination also modulate diverse processes such as adaptation, speciation, and introgression (Samuk et al. 2017; Dapper and Payseur 2017; Stapley et al. 2017; Schumber et al. 2018). There is also a growing appreciation that recombination rate is itself a trait that varies and evolves (Dumont and Payseur 2008; Hunter et al. 2016; Johnston et al. 2016; Ritz et al. 2017; Stapley et al. 2017; Samuk et al. 2020). Accordingly, there has been great interest in efficient and accurate methods for estimating recombination rates.

Current methods for estimating recombination rates fall into 2 broad classes of methods: direct and indirect (Peñalba and Wolf 2020). Of the direct measures, the 3 most popular approaches are linkage mapping, gamete sequencing, and cytological methods. With classical linkage mapping, map distances between genetic markers are measured by quantifying recombinant markers in the context of a genetic cross or pedigree (Broman 2010; Rastas 2017). The resolution of this approach is limited only by marker density and the sample size of individuals, but larger sample sizes can be grueling to achieve in the laboratory or unavailable in some populations. Furthermore, identifying suitable diagnostic mapping markers can be limiting in some cases (e.g. in a highly homozygous population; Broman 2010). Direct sequencing of pools of recombinant gamete genomes from single individuals using long/linked read sequencing is a newer approach that alleviates many of the issues of traditional mapping, but still requires differentiated markers to score crossover events between homologous chromosomes (Dréau et al. 2019; Rommel Fuentes et al. 2020; Xu et al. 2020). Cytological methods bypass this requirement by directly visualizing recombination-associated protein complexes in cell populations undergoing meiosis (Peterson et al. 2019; Peterson and Payseur 2021). However, the cytological methods are limited by the spatial resolution at which such visualization can occur (e.g. the resolution of immunostained gamete karyotypes; Peterson et al. 2019).

Because all direct methods of measuring recombination rates are fairly laborious, there has been increased interest in indirect measures of recombination rate that leverage readily available population genetic data. Chief among these are model-based methods that infer rates of recombination from patterns of linkage disequilibrium (LD; Auton and McVean 2007; Chan et al. 2012; Kamml et al. 2016; Spence and Song 2019). These methods attempt to estimate recombination rates by statistically fitting recombination rates (derived from population genetic models/simulations) to observed...
patterns of LD. Rather than inferring recombination rate directly, LD-based estimators infer a population scaled recombination rate, \( \rho = 4N_e r \), where \( N_e \) is the effective population size and \( r \) is the theoretical per-generation recombination rate. LD-based methods are attractive because they (1) generally only require population-scale genomic data and (2) are very fast, often only requiring several computational hours or less (Spence and Song 2019) and (3) are informative of time-averaged population historical recombination rates (McVean and Auton 2007). Accordingly, LD-based estimates of recombination rates have become extremely popular, and now vastly outnumber direct measures in the literature (Stapley et al. 2017; Peñaiba and Wolf 2020). These methods have also begun to be used to perform interpopulation comparisons of recombination rates (Stapley et al. 2017; Peñaiba and Wolf 2020).

Like all models, LD-based estimators of recombination rate make a variety of simplifying assumptions about the populations of interest. For one, they generally assume that the populations/loci of interest are evolving largely neutrally and have reached population genetic equilibrium in a number of ways (Stumpf and McVean 2003). In particular, most methods assume that the populations being studied have reached an equilibrium between recombination and population scaled mutation, such that LD accurately reflects patterns of recombination rate (McVean 2007). Furthermore, it is generally assumed that any form of selection that might distort patterns of LD (e.g. sweeps) has not recently occurred (Chan et al. 2017; Peñaiba and Wolf 2020). Finally, these methods make the general assumption that demographic processes that distort genome-wide patterns of LD, such as population size changes, have not occurred (recall that \( p \) is directly dependent on \( N_e \), Auton and McVean 2007).

While some of these assumptions may be robust to violation, work has shown that some violations can result in biased estimates. For example, Dapper and Paysseur (2018) showed that recombination estimates from LDhat (McVean and Auton 2007) are highly sensitive to changes in population size. This can be alleviated in some cases by incorporating known changes in population size into the estimation procedure, such as implemented in the software pyrho (Spence and Song 2019).

Along with changes in population size and selection, another process that can greatly alter patterns of LD is gene flow. Gene flow and subsequent admixture between diverged populations can have complex effects on patterns of LD within each population (Nei and Li 1973; Ohta 1982). These effects range from large and genomically variable increases in LD due to segregation of divergent haplotypes, to genome-wide decreases in LD as populations become coupled and increase local \( N_e \) (Nei and Li 1973; Ohta 1982). While this is now widely accepted that gene flow is commonplace in natural populations (Barton 2001; Mallet 2005; Waples and Gaggiotti 2006; Suvorov et al. 2022), there has not been a systematic study of the effects of gene flow on LD-based measures of recombination. Furthermore, it remains unclear how gene flow (or any other violation of assumptions) impacts our ability to detect differences in recombination rate between (as opposed to within) populations using LD-based methods.

Here, we address these issues using forward-time population genetic simulations. We attempt to answer 2 specific questions. First, how does gene flow between populations affect the precision and accuracy of LD-based estimates of recombination rate within populations? Secondly, how does gene flow affect our ability to detect evolved differences in recombination rate between populations? Our primary goal is to answer these questions in the context of a core set of realistic demographic scenarios, and not perform an exhaustive exploration of parameter space. Overall, we hope to help investigators understand key sources of bias in LD-based estimates of recombination rate in natural populations and highlight areas of future development.

**Methods**

**Code availability**

All scripts used in the analyses described below are available as a repository on Github (http://github.com/ksamuk/LD_recomb).

**Forward time simulations with SLiM**

To explore how the timing and amount of gene flow affect estimates of recombination rate, we performed forward-time population genetic simulations using SLiM version 3.3 (Haller and Messer 2019). The basic form of all the simulations was an isolation-with-migration scenario: a single ancestral population diverges into 2 subpopulations with a static amount of bidirectional gene flow (Fig. 1). Populations were composed of diploid individuals with 100-kb genomes arranged in a single chromosome. We used genome-wide average estimates of effective population size, mutation rate, and empirical recombination rate from natural populations of Drosophila melanogaster (Adrion, Cole et al. 2020). Per site mutation rate = 5.49 \times 10^{-5} (Li and Stephan 2006); per site recombination rate = 2.23 \times 10^{-6}, (average of chromosome 2R, Comeron et al. 2012); \( N_e = 1.72M \) (Li and Stephan 2006). Recombination and mutation rates were conservedly modeled as uniform across the 100-kb genome. Following standard practice for forward-time simulations, all simulations were run with an in silico population size of \( N = 1,000 \), and simulated mutation and recombination rates scaled by a factor of \( N_e/N \) as per the SLiM manual (Haller and Messer 2019). Note that generation times are also subject to scaling, and for simplicity, we will refer to all generations in terms of back-transformed actual generations rather than SLiM generations (1 SLiM generation \( \approx 1,751 \) actual generations with our scaling factor).

**Parameter space**

To explore how variation in gene flow affects estimates of recombination, we varied the amount of gene flow over 5 orders of magnitude: 0, 0.01, 0.1, 1, 10, 100, in standard units of \( 4N_e m \) (the product of the effective population size and the migration rate). These values were chosen to encompass total isolation (\( N_e m = 0 \)), limited gene flow (\( N_e m = 0.01-0.1 \)), moderate gene flow in interconnected metapopulations (\( N_e m = 1-10 \); Morjan and Rieseberg 2004; Waples and Gaggiotti 2006), and a scenario of a nascent hybrid swarm (\( N_e m = 100 \)). We also varied the timing of the onset of gene flow, with gene flow beginning either immediately after divergence or after a period of isolation. We performed preliminary simulations to determine a period of isolation (~1.7M generations in our case) that produced levels of genomic divergence (Supplementary Fig. 1) similar to those observed in natural population pairs that exhibit genome-wide genetic divergence but still actively exchange genes (~FST ~ 0.4; Morjan and Rieseberg 2004; Roux et al. 2016). Finally, to explore how gene flow impacts the detection of population differences in recombination rate, we modeled scenarios where recombination rate either remains constant in both subpopulations or instantaneously increases by a factor of 2 at the time of divergence in one of the 2 subpopulations (always subpopulation 2). This magnitude of this difference is well within the range of variation in recombination rate reported for a wide variety of species (Stapley et al. 2017). In biological terms, an instantaneous increase in population recombination rate could be readily mediated by an environmental change (e.g. temperature, Lloyd et al. 2018), a change in mating system (Brandvain and Wright 2016), or whole-genome duplication (Tiley and Burleigh 2015). We note that this instantaneous change
Details of demographic events

Each simulation began with a single population of size $N_{e,anc}$, which evolved for a 35M generation burn-in period (following the general practice of a 10-20 $N_e$ burn-in period; Haller and Messer 2019). This initial period was followed by divergence into 2 subpopulations, each with size $N_{e,anc}/2$. Gene flow (for cases where $N_{em} > 0$) began immediately at the time of divergence or after a 1.7M generation period of isolation and was symmetrical in magnitude and bidirectional. Changes in recombination rate occurred at the time of divergence and instantaneously applied to all individuals in subpopulation 2 only.

Starting at the time of divergence and thereafter in intervals of 1,751 generations, we collected a random sample of 25 individuals (a total of 50 haploid genomes) from each population and saved their complete genotypic at all sites in VCF format. We stopped the simulations after 51,000 generations. Each parameter combination was replicated 100 times, for a total number of $\sim 48,000$ population samples.

Estimation of recombination rate using pyrho

While there are a variety of LD-based estimators of recombination rate, we elected to use pyrho (Spence and Song 2019) for estimation in this study. It shares its statistical foundation with the most widely used LD-based estimators (LD-hat and LD-helmet; McVean and Auton 2007; Chan et al. 2012) while also having the ability to account for changes in effective population size such as we are modeling here (Spence and Song 2019). As such, any estimation biases caused by gene flow will likely affect those approaches as least as much as they affect pyrho. Direct comparisons with other methods are complicated by the fact that pyrho is the only model-based method that adequately accounts for changes in effective population (Adrion, Galloway et al. 2020).

We followed the recommended practices for inferring recombination rate using pyrho (https://github.com/popgenmethods/pyrho). We parameterized the initial lookup tables using the effective population size and mutation rates used in the simulations (unscaled in this case). To account for changes in effective population size, we created lookup tables that accounted for a change of $N_{e}/2$ (1.72M to 8.6M) in time steps of 1,751 generations in the past. This allowed us to have an appropriately timed lookup table for each step of the simulation. We used the built-in methods to infer the hyperparameters of window size (best fit 100) and block penalty (best fit 1,000). Using this baseline, we inferred recombination rates using the genotype data (VCF format) from both subpopulations at each time point, for a total of $\sim 96,000$ pyrho fits. All computations were performed using the Duke University Computing Cluster, running CentOS Version 8.

Statistical analyses

We performed all data processing and visualization using the tools of the tidyverse package in R 4.0.3 (Wickham 2017; R Core Team 2018). To examine how gene flow between populations affects the accuracy of LD-based estimates of recombination, and the context of the various factors explored in our simulations, we performed an analysis of variance using a linear mixed model with Gaussian errors fitted via the lmer() function from the lme4 package (Bates et al. 2015). This model had the following form: Recombination rate = (1|simulation replicate) + (1|simulation generation) + gene flow magnitude + recombination rate change, where (1|factor) denotes a random intercept. All variables were standardized (mean-centered and scaled by standard deviation) prior to analysis. To simplify interpretation, we fitted separate models for the continuous gene flow and secondary contact scenarios.

Results

Inference when recombination rate is identical between populations

When the recombination rate remained constant between diverging populations, we found that gene flow introduced 2 types of systematic biases in estimates of recombination rate within populations (Fig. 2a). These effects began when $N_{em} \geq 1$ in both the
continuous gene flow and secondary contact models. First, in the model of continuous gene flow, when \( N_{em} \geq 1 \), we observed a systematic increase (overestimate) in estimated rates of recombination in both populations (Fig. 2, a and b, top row, \( N_{em} = 1–100 \)). This increase was statistically significant [Type III Wald chi-square = 5090.07, \( P < 2.2 \times 10^{-16} \); coefficient for gene flow = 0.63–0.67 (95% CI), \( t(19495) = 71.34, P < 0.001 \)]. When the migration rate was moderate to high (\( N_{em} = 10–100 \)), the recombination rate was overestimated by \( \sim 10–20\% \) (Fig. 2b). This effect is consistent with migration causing the populations to become coupled, behaving as a single population with a larger \( N_e \) and thus inflating the population-scaled estimate of recombination rate.

In contrast to the continuous gene flow case, under a model of secondary contact, there was a marked systematic decrease (underestimate) of recombination rates, which also became visible when \( N_{em} \geq 1 \) (Fig. 2, a and b, bottom row, \( N_{em} = 1–100 \)). This decrease was statistically significant [Type III Wald chi-square = 1512, \( P < 2.2 \times 10^{-16} \); coefficient for gene flow = \(-0.54–0.49 \) (95% CI), \( t(31846) = -38.88, P < 0.001 \)]. The magnitude of this decrease was substantial: on average, populations experiencing \( N_{em} = 1 \) had recombination rates about 20% lower than expected, with this increasing to 50% when \( N_{em} = 10 \) or higher (Fig. 2, a and b). This decrease was accompanied by a statistically significant increase in the variance of recombination rate estimates, especially for \( N_{em} = 1–10 \) compared to \( N_{em} < 1 \) (Fig. 2a, bottom row; \( F \)-test for equivalency of variance, \( F(10,429,13,860) = 0.20863, P < 2.2 \times 10^{-16} \)). A systematic increase in the mean and variance of LD within populations is consistent with...
allele frequency differences between populations manifesting as migration-associated LD, and deflating estimates of recombination rate. When gene flow was very high, there was a visible recovery of estimated recombination rates (Fig. 2a, bottom row, N_em = 100), presumably due to migration homogenizing allele frequencies and increased effective population sizes increasing the rate at which recombination breaks down migration-associated LD.

When comparing recombination rates between p1 and p2, the “coupling” bias observed in the continuous migration scenario did not appear to systematically affect the ratio of recombination rate between the 2 populations (Fig. 2c, Continuous Migration). However, in keeping with the previous result, migration-associated LD in the secondary contact model appeared to greatly increase the between replicate variance in the ratio of recombination rates between populations when N_em ≥ 1 (Fig. 2c, Secondary Contact).

**Inference when recombination rate differs between populations**

When recombination rates diverged between populations, we also observed the 2 forms of bias described above (Fig. 3). The estimates from the continuous gene flow scenario exhibited a statistically significant increase [Type III Wald chi-square = 8,936.44, P < 2.2 × 10^-16; coefficient for gene flow = 0.65–0.67 (95% CI), t(19495) = 94.53, P < 0.001] whereas estimates from the secondary contact model exhibited a statistically significant decrease [Type III Wald chi-square = 1,512, P < 2.0 × 10^-16; coefficient for gene flow = –0.27–0.22 (95% CI), t(34505) = –23.22, P < 0.001]. However, the results differed from simulations with constant recombination rates in a number of important ways. First, there was a clear difference between the continuous migration and secondary contact models in the overall trajectory in the population-specific estimates of recombination rate (Fig 3a). In the continuous gene flow models, there was an overall positive trend for the estimates of recombination rate in p2 even in the absence of gene flow (Fig 3a, continuous migration). This was presumably caused by a lag in the establishment of equilibrium levels of LD within p2 that reflect the new recombination rate (which spontaneously changed at the time of divergence). This lag resulted in the recombination rate in p2 being consistently underestimated (because it had not reached its new equilibrium), in addition to the coupling effect observed previously (Fig 3, b and c, continuous migration). In the case of the secondary contact model, we did not observe the same positive trend for recombination rate estimates in p2, likely because the isolation period (1.7M generations) was sufficiently long enough for p2 to establish an equilibrium level of LD prior to secondary contact (Fig 3a, Secondary Contact).

**Additional simulations**

To examine the robustness of our results, we explored 2 additional demographic scenarios. First, we repeated our simulations using an effective population size of 1,720 (1/0.001th of the D. melanogaster Ne). These simulations produced broadly similar results, with all biases we identified using a larger Ne also appearing in the presence of a smaller Ne (Supplementary Figs. 2 and 3). Furthermore, there was an apparent increase in variance of recombination estimates in many cases, suggesting that the issues we identified may be considerably worse in species with smaller effective population sizes (Supplementary Fig. 2c).

Finally, using these smaller effective population size simulations as a base, we also explored how asymmetry in gene flow may affect the estimation of recombination rate. To do this, we performed simulations in which gene flow is unidirectional from p2 into p1. Asymmetrical gene flow resulted in an increase in both the bias and variance of recombination rates estimates, with the worst effects again manifesting at moderate levels of gene flow (Supplementary Figs. 4 and 5). As such, it appears that our core simulations (large Ne, symmetrical gene flow) may represent a “best case scenario” for mitigating biases, and departures from this balanced scenario appear to only worsen the prospects for accurate population genetic estimation of recombination rates.

In keeping with the scenario with constant recombination rates, starting at N_em ~ 1, migration-associated LD resulted in the systematic underestimation and increase in variance for estimated recombination rates within both p1 and p2 [Fig. 3b, Secondary Contact; Type III Wald chi-square = 538.97, P < 2.0 × 10^-16; coefficient for gene flow = –0.27–22 (95% CI), t(34505) = –23.22, P < 0.001; F-test for equivalency of variance, F(7,734, 13,579) = 0.2174, P < 2.2 × 10^-16]. In addition, the observed divergence in recombination rate between p2 and p1 (which was always expected to be +2 cm/Mb) decreased with increasing levels of gene flow (Fig. 3, b and c; Secondary Contact). This effect would likely result in an increase in false negatives with increasing gene flow (i.e. finding no difference in recombination rate between populations when there is in fact one). This decrease in the observed divergence between populations is again likely the outcome of the population-specific levels of LD becoming coupled/merged at moderate to high levels of gene flow, resulting in the populations exhibiting LD (and hence recombination rate estimates) intermediate to what would be expected in the absence of gene flow.

**Discussion**

Accurate estimates of recombination rate are key to understanding the causes and consequences of recombination rate variation in natural populations. With the increasing availability of genome-wide sequencing data, LD-based estimators of recombination rate have become widely used in a large variety of taxa. However, while gene flow is widely known to shape patterns of LD in populations, the effect of gene flow on LD-based estimators of recombination rate remains largely unexplored. Here, we used forward-time simulations to show that 1) moderate to strong gene flow can introduce substantial bias into LD-based estimates of genome-wide recombination rate and 2) the nature of this bias depends on the demographic and evolutionary history of the populations in question.

Our results here are consistent with theoretical predictions that gene flow between populations can affect LD: increasing in the magnitude and variance of LD at low migration rates as well as reducing LD via the “coupling effect” we observed at higher rates of gene flow. Our study shows how these predictions play out with modern methods and genomic data, and also provides a sense of the magnitude of the potential degree of misestimation—in our case, ranging from 20 to 50 percentage points in cases of moderate gene flow. For comparison, a recent study of population-level differences in recombination rate in Drosophila pseudoobscura revealed genetically based interpopulation differences on the magnitude of ~10% measured using replicated linkage maps in each population (Sarmuk et al. 2020). Using LD-based estimators, an observed difference of this magnitude could be spuriously generated by modest levels of gene flow alone, or missed altogether due to coupling at higher levels of gene flow.
In addition, the specific magnitude and direction of the bias introduced by gene flow are difficult to know without precise knowledge of the population/demographic histories of the populations in question. This should give pause to anyone planning on using LD-based methods to infer recombination rate in nonequilibrium populations.

One key question is whether there are methods to control for or counteract the increased variance and/or biases in the estimation of recombination rate caused by gene flow. One approach could be to identify and remove introgressed haplotypes from datasets prior to inferring recombination rate, thereby removing migration-associated LD. This would require “pure” samples from the source populations, such that the population of origin could be assigned to haplotype blocks (Dias-Alves et al. 2018). However, this method would require gene flow to be low enough that coupling (of both LD and allele frequencies) has not occurred. The upward bias and increased variance in recombination rate that occurs as a result of coupling, together with the homogenization of allelic differences between populations at higher levels of gene flow will likely make a “filtering” scheme very difficult (perhaps impossible) to achieve. One other approach may be to attempt to jointly estimate a demographic model along with population-specific recombination rates, as has been done with mutation rates (DeWitt et al. 2021). However, given the existing complexity and uncertainty in inferring demographic models, we suspect it may be difficult to disentangle the complex interdependencies between gene flow, population size, and estimates of recombination rate.

Fig. 3. The relationship between inferred recombination rate and the migration rate in simulated populations where recombination rate increases by a factor of 2 in one subpopulation. (a) Inferred recombination rates for individual simulations at varying levels of migration. Each plot shows inferred rates for simulation replicates (transparent lines) of population 1 (red, unchanged recombination) and population 2 (blue, increased recombination) for a single migration rate. Dashed lines show the expected inferred value in the absence of gene flow (inferred from \( N, m = 0 \)). (b) Summarized inferred recombination rates (y-axis) for each level of migration (x-axis) from the simulations in a. Points are mean values and error bars depict standard deviations (summarized across all generations). Dashed lines show the expected inferred value in the absence of gene flow for each population (i.e. the mean value for \( N, m = 0 \)). (c) The inferred difference in recombination rate between population 1 and population 2 (\( p_2-p_1 \)) as a function of migration rate. Points and error bars are as in b.
On a related note, in our simulations we had perfect knowledge of the demographic histories of both populations (ancestral and derived population sizes; divergence time), which was used to parameterize the correction procedure employed by pyrho. In the vast major of empirical cases, demographic history would need to be separately estimated prior to parameterizing pyrho. Such demographic inference is itself error prone and subject to a wide variety of potential biases (Marchi et al. 2021) and these errors would propagate into estimates of rho (see Dapper and Payseur, 2018). Furthermore, the interactions between selection, gene flow, and recombination likely further complicate inference of rho. For example, many studies have now shown a negative correlation between recombination rate and introgression mediated by alleles causing reproductive isolation (Aeschbacher et al. 2017, Samuk et al. 2017, Schumer et al. 2018). This suggests that biases in rho estimation introduced by gene flow could themselves vary with genomic context. As such, the biases and error rates identified here represent a “best case” scenario, and would be in addition to any errors due to misestimation of the demographic history or the effects of reproductive isolation.

Together with previous work (Dapper and Payseur 2018), our results suggest that LD-based estimates of recombination rate need to be interpreted with great caution when studying nonequilibrium populations. Indeed, these methods are likely only appropriate when populations can be assumed to be evolving in the long run under the assumptions of demographic equilibrium more readily (Ochoa and Storey 2019; Samuk et al. 2021). In a review of population comparisons of traits by Leinonen et al. (2008), such populations comprise around 20–30% of the cases identified. It is also worth noting that it is not the case that 2 populations being studied have to be exchanging genes themselves (e.g. which would not be the case when studying 2 reproductively different species), but just that one or more of the populations are exchanging genes with some other population (e.g. an unsampled population of the same species).

If many LD-based estimates are incorrect, why do published LD-based estimates of recombination rate correlate well with direct estimates, e.g. from genetic maps? (McVean and Auton 2007; Chan et al. 2012, Smukowski Heil et al. 2015). There are several considerations. First, the correlations that have been reported are by no means perfect (e.g. ~Spearman’s Rho of 0.6: Smukowski Heil et al. 2015; $r^2 = 0.37–63$: (Chan et al. 2012) and depend greatly on the genomic scale at which they are measured (Smukowski Heil et al. 2015). Second, simple correlations between LD-based and empirical estimates do not speak to genome-wide differences in the estimates of recombination rate, such as those due to the coupling effects we observed. Such effects would be visible as differences in the intercept of a linear regression, rather than the $R^2$, for example. Finally, the species where these correlations have been examined (humans and D. melanogaster) may meet the assumptions of demographic equilibrium more readily (Ochoa and Storey 2019; Suvorov et al. 2022). While such assumptions may be reasonable for these populations, for which LD-based estimators were originally developed, they are much less likely to hold in many natural populations. Notably, they are likely rarely met in populations that have recently adaptively diverged in the presence of gene flow, which have lately been the subject of increased research interest (Ravinet et al. 2017; Linck and Battey 2019). The equilibrium assumption is also likely not valid in populations in which the recombination rate has recently changed (Brandvain and Wright 2016), reducing the utility of these estimates for studying the rapid evolution of recombination rates.

While we only focused on a single implementation of one type of LD-based estimator of recombination (pyrho), it is likely that other population genetic methods will also suffer from the effects we describe here. LD is the “information” used by all estimators, either directly as in methods like LDjump (Herrmann et al. 2019) or indirectly as in machine learning methods like ReLERNN (Adrion, Galloway et al. 2020). That said, in the case of the latter method, it may be possible to overcome some of the issues we have identified if the training datasets were simulated with an accurate demographic model. As such, the distorting effects of gene flow on LD need to be carefully considered when applying any statistical methods for inferring recombination rate approaches. We also stress that our simulations do not suggest that LD-based estimators and their implementations are wrong per se, but rather that the assumptions under which LD-based estimates are biologically accurate are readily violated by levels of gene flow and divergence common seen in natural populations.

Finally, the biases we have identified likely affect the identification of recombination cold/hotspots and the “landscape” of recombination in general. For example, if introgression is itself variable across the genome, this could result in the biases we have identified here (1) covarying with introgression and (2) creating false heterogeneity in recombination estimates. The increase in variance we identified could also result in (apparent) increased heterogeneity in recombination across the genome. In terms of identifying hotspots, the difference in recombination rate between hot and cold spots in most species vastly exceeds the 20–50% differences we described here, and thus the biases we identified may not be an issue for the identification of extreme hotspots per se.

**Conclusion**

Studying variation in recombination rate is difficult. LD-based methods for inferring recombination rate are attractive in their data requirements but require strong assumptions to be met. As we have shown here, gene flow readily violates these assumptions and introduces biases and decreases in precision, in a variety of ways that are difficult to identify in a given study population. This is problematic because gene flow is extremely common in natural populations. How should we proceed? Rather than attempt to squeeze blood from the proverbial stone, we believe that the most straightforward solution to the problems we outline here is simply to prioritize the use of direct, empirical methods for measuring of recombination rate. This decision is made hopefully simpler with the increased ease and low cost of creating traditional linkage maps and performing gamete sequencing. That said, LD-based approaches remain important tools for hypothesis generation, and when paired with direct estimates of recombination rate can provide a detailed picture of both the past and present landscape of recombination rates in natural populations.
Data availability
All codes used to generate the simulated data used in this study are available as a repository on Github (https://github.com/ksa-muk/LD_recomb).
Supplemental material is available at G3 online.

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Conflicts of interest
None declared.

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