

# The short-term, genome-wide effects of indirect selection deserve study: a response to Charlesworth and Jensen (2022)

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**Running Title:** Response to Charlesworth and Jensen

We recently published a paper quantifying the genome-wide consequences of natural selection, including the effects of indirect selection due to the correlation of genetic regions (neutral or selected) with directly selected regions (Gompert *et al.*, 2022). In their critique of our paper, Charlesworth & Jensen (2022) make two main points: (i) indirect selection is equivalent to hitchhiking and thus well documented (i.e., our results are not novel), and (ii) that we do not demonstrate the source of linkage disequilibrium (LD) between SNPs and the *Mel-Stripe* locus in the *Timema cristinae* experiment we analyze. As we discuss in detail below, neither of these are substantial criticisms of our work.

First, indirect selection and hitchhiking are related but not equivalent concepts. Our focus on the short-term consequences of long-range LD (i.e., across the genome and not restricted to tightly-linked regions) sets our work apart from most population genetic studies of hitchhiking. In this context, we take the opportunity here to expand on why readers of *Molecular Ecology* might be interested in the short-term consequences of indirect selection.

Second, the consequences of LD for indirect selection that our study focused on hold regardless of the source(s) of LD. Drift and local migration, as well as natural selection itself, can all create a degree of long-range LD between selected sites and neutral unlinked variants, particularly in small, finite populations (Fig. 1). We demonstrated the existence of this long-range LD with respect to the *Mel-Stripe* locus in *Timema cristinae* (which controls color), as well as in two other cases involving the *Agouti* gene and coat color in deer mice (*Peromyscus maniculatus*) and the *Ectodysplasin* gene and body armor in stickleback fish (*Gasterosteus aculeatus*). Although this long-range LD is transient compared to physically linked sites, our point was that selection on a target locus can deterministically affect the direction of allelic change at even unlinked neutral sites for short periods of time (a few generations), before being dissipated by recombination and random assortment. Moreover, new long-range associations are constantly reforming between selected and other sites in finite populations due, in part, to the sources creating LD discussed by Charlesworth & Jensen (2022). In a metaphoric sense, selected sites will be constantly picking up new unlinked passengers in

the genome and taking them for short rides. Thus, while we appreciate the points raised by Charlesworth & Jensen (2022), they do not detract from our main thesis. We elaborate on these misunderstandings below in hopes that it clarifies the role and significance of indirect selection in evolution for the readers of *Molecular Ecology*.

The concept of indirect selection was introduced by Pearson (1903) over 70 years before the term hitchhiking was first used to describe the within-generation change in the distribution of phenotypes for traits not directly affecting fitness but instead that were correlated with traits directly under selection. Such correlations might arise because of LD between loci affecting different traits, pleiotropy, or a common effect of the environment on sets of traits. Thus, in contrast to standard usage of hitchhiking (discussed more in the next paragraph), indirect selection does not emphasize physical linkage nor does it focus on the long-term consequences of selection on patterns of molecular variation. The concept of indirect selection was then brought to the widespread attention of evolutionary biologists in the 1980s, for example, by Lande & Arnold (1983) introducing the use of multiple regression to disentangle direct versus indirect selection on traits. Since this time, thousands of studies have applied the methods of Lande & Arnold (1983) to measure indirect selection (reviewed in Kingsolver *et al.*, 2001, 2012), with recent applications to measuring indirect selection on genetic loci (e.g., Gompert *et al.*, 2014; Thurman & Barrett, 2016; Gompert *et al.*, 2017; Exposito-Alonso *et al.*, 2019).

The concept of hitchhiking was introduced by Maynard Smith & Haigh (1974) to describe the increase in frequency of an allele present on a chromosome where a new beneficial mutation arises that is caused by LD between the allele and the beneficial mutation. This concept emphasizes physical linkage and the evolutionary consequences of new mutations, especially in the context of genetic diversity. As such, hitchhiking can be viewed as a consequence of indirect selection, where physical linkage is involved. Hitchhiking was extended to the case of linked, indirect selection against deleterious alleles, that is background selection by Charlesworth *et al.* (1993). More recent work has investigated soft or partial selective

sweeps and hitchhiking caused by selection on standing genetic variation (e.g., Hermisson & Pennings, 2005; Prezeworski *et al.*, 2005; Barrett & Schluter, 2008; Hermisson & Pennings, 2017). Thus, while indirect selection and hitchhiking are clearly closely related terms, they have distinct historical usage and the distinction between them is more than semantic. We here argue that indirect selection is best used to describe the immediate indirect effects of selection on one trait or gene caused by correlations with fitness (or other selected traits or genes), whereas hitchhiking best describes the longer-term consequences of this for patterns of molecular variation especially when physical linkage is involved. With that said, the recent (alternative) definition of hitchhiking given by Charlesworth & Jensen (2021) based on the additive genetic covariance between a trait (or gene) and fitness via the Price-Robertson equation (Robertson, 1968; Price *et al.*, 1970) is in essence equivalent to our treatment of indirect selection and the way in which this term has been mostly used in evolutionary biology since Pearson (1903).

We think that this (admittedly subtle) distinction between indirect selection and hitchhiking helps to clarify the novelty of our recent work (Gompert *et al.*, 2022). As noted by Charlesworth & Jensen (2022), and as made clear in our original manuscript (Gompert *et al.*, 2022), the long-term consequences of indirect selection, and specifically of hitchhiking (including background selection) on patterns of genetic variation are well established, with especially compelling evidence for the impact of background selection on patterns of diversity in many organisms (e.g., Begun & Aquadro, 1992; Comeron, 2014; Charlesworth & Campos, 2014; Pouyet *et al.*, 2018). Also as noted by Charlesworth & Jensen (2022), this of course implies that indirect selection must operate on shorter timescales. Despite this theoretical truism, the importance of indirect selection for short-term evolutionary or ecological dynamics has received far less empirical attention. Specifically, this cannot be determined from static studies of molecular variation alone but instead requires studies of contemporary evolution in natural populations, ideally combined with field and lab experiments that measure selection and its immediate consequences across the genome.

In Gompert *et al.* (2022) we aimed to begin to fill this knowledge gap and provide evidence that indirect selection has the potential to have measurable short-term impacts on evolution. These impacts include indirect selection on genetic variants not physically linked to a focal directly selected locus. Although we do focus on the color locus *Mel-Stripe* in *T. cristinae*, the only system criticized by Charlesworth & Jensen (2022), we stress that the conclusions noted above rest on the collective analysis of data sets in stick-insects, stickleback and mice. Moreover, the existence (though not quantitative extent) of genome-wide indirect selection is a mathematical certainty in any finite population experiencing selection. In the stick-insect example, we showed that LD (measured by  $r^2$ ) for 3% of all SNPs not on the same chromosome as *Mel-Stripe* was  $> 0.01$  and that this exceeded 0.10 for 64 SNPs. Importantly and also distinct from most work on hitchhiking, we extended the results in stick insects to show that LD with numerous (unknown) causal variants likely affecting fitness causes even more genomically widespread indirect selection (i.e., due to polygenic selection, see Gompert *et al.*, 2022 for details).

Should we invest effort in investigating possible short-term (i.e., a one or several generations) long-distance (i.e., among distant loci including unlinked loci) effects of indirect selection? As pointed out both by Charlesworth & Jensen (2022) and ourselves (Gompert *et al.*, 2022), recombination and independent assortment cause LD to decay over time. Thus, long-distance LD is not expected to persist for long periods of time. Nonetheless, the short-term, long-distance effects of indirect selection should be considered for several core reasons. First, incorporating the short-term effects of indirect selection provides a more mathematically precise and conceptually appropriate model of evolution. For example, with indirect selection, the expected single generation change in allele frequency for a neutral locus in LD with loci affecting fitness is not 0 (i.e.,  $E[\Delta p] \neq 0$ ), as it would be in standard models of genetic drift (Wright, 1931). This difference might be slight in most cases, but still it is real, even if past drift was the cause of LD. Such indirect selection might be especially important in field or lab studies of strong selection in small populations. Second, in some cases, the

short-term effects of indirect selection could have meaningful ecological consequences. Even for physically unlinked loci, LD can persist for several generations (it decays by half each generation), and there is now compelling evidence that even short-term evolutionary change, especially in fluctuating or heterogeneous environments, can have ecological consequences for entire communities (reviewed in Hendry, 2020). Third, some species have long generation times (e.g., many tree species), and thus patterns of LD might persist for hundreds or thousands of years in absolute time (i.e., the entire modern period of human-induced climate change). Thus, although further work is required to compare the effects of indirect selection across timescales and genomic regions, we do not think there is sufficient evidence at present to completely dismiss the study of short-term, long-distance effects.

The other major critique raised by Charlesworth & Jensen (2022) was that we did not determine the source of LD in the *T. cristinae* experiment. As noted above, this was because the consequences of LD for indirect selection, the focus of our study, are not conditional on the causes of LD in the population. Thus, this is not a valid criticism of our study. Still, we do agree that this is an interesting question worth considering and we take the opportunity to do so here. We agree that each of the six mechanisms identified by Charlesworth & Jensen (2022) could contribute to the observed LD. We think most of these are unlikely, but comment on all briefly here, before discussing what we think are the most likely explanations in more detail. The first possibility raised was that SNPs in LD with *Mel-Stripe* were in fact physically linked to *Mel-Stripe* but were spuriously placed on other chromosomes. The overall quality of our reference genome (mostly large scaffolds derived from a Chicago genomic library) and linkage groups (constructed from crosses) makes this unlikely to be the primary cause of LD (Nosil *et al.*, 2018) (more recent, but as of yet unpublished chromosome-level genome assemblies further support our conclusion). The second possibility raised is genetic drift, which Charlesworth & Jensen (2022) dismiss but we think is a highly probable mechanism as we discuss below. Their third mechanism is very recent admixture. While this is unlikely in light of past genetic analyses in *T. cristinae* (which were not considered by Charlesworth &

Jensen, 2022) (e.g., Nosil *et al.*, 2012; Soria-Carrasco *et al.*, 2014; Riesch *et al.*, 2017), we do think that population structure and ongoing gene flow (combined with selection) contribute to the observed LD. The fourth mechanism was a recent bottleneck, which is possible but not supported by observations in the field over the last 29 years, where population size of the source population fluctuated but has not dropped extensively (e.g., Nosil *et al.*, 2018 for a long-term study). Fifth, Charlesworth & Jensen (2022) suggested but mostly dismissed an ongoing selective sweep at *Mel-Stripe*; we agree that this is unlikely given evidence for long-term balancing selection on this locus (Lindtke *et al.*, 2017; Nosil *et al.*, 2018; Villoutreix *et al.*, 2020). Lastly, Charlesworth & Jensen (2022) suggest a possible role for epistatic interactions for fitness between *Mel-Stripe* and other genetic variants genome wide. We agree with them that this is unlikely to be a major mechanism (i.e., it could apply but more so to one or a few loci).

Thus, in terms of likely explanations for the observed LD, we think genetic drift and gene flow warrant prime consideration. In their critique, Charlesworth & Jensen (2022) are mostly dismissive of genetic drift because of the size of the source *T. cristinae* population from which the experimental population was sampled ( $\sim 1000$  individuals). However, their criticism fails to distinguish between census and effective population size. Past work in *T. cristinae* suggests that  $N_e$  is considerably smaller than the census population size, and probably on the order of 10% of the census size (Soria-Carrasco *et al.*, 2014; Nosil *et al.*, 2018), consistent with broad patterns in other organisms (Frankham, 1995). Simple simulations show that with  $N_e \sim 100$ –200, drift could readily create the patterns observed (Fig. 2). Second, gene flow likely contributes to the observed patterns of LD. Past work, including demographic modeling from population genetic data suggest gene flow at least at smaller spatial scales, including for differentiated regions under divergent selection where gene flow will create LD (Nosil & Crespi, 2004; Nosil *et al.*, 2006, 2012, 2018). However, we re-iterate that the causes of LD have little bearing on the core conclusions of our original study (Gompert *et al.*, 2022).

We want to conclude by thanking Charlesworth & Jensen (2022) for helping us to clarify the issues discussed above. We think that our disagreements with them mostly reflect differences in emphasis between studies of indirect selection (prominent in field studies of natural selection) and hitchhiking (prominent in population genetics and molecular evolution). We hope that this discussion provides cross-talk between these important sub-disciplines. We fully agree with Charlesworth & Jensen (2022) that the long-term consequences of hitchhiking, including background selection, are well known and that we are now at the point where background selection should be part of our standard null models in molecular evolution (Comeron, 2017). We also concur that resolving the processes creating long-range LD in *Timema* and other systems is important. However, it is the subsequent effects that selection may have indirectly on unlinked sites due to the existence of long-range LD, as demonstrated in the manipulative transplant experiment for *T. cristinae*, that we highlight as the take-home message of our paper. Thus, we stand by our original argument that the short-term, long-distance consequences of indirect selection on genetic loci deserve more attention. Ultimately, it will be further empirical work that reveals the relevance of this process for ecology and evolution, particularly for finite populations experiencing polygenic and varying selection on many traits.

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## 258 **Data availability**

259 No new data were analyzed in this paper, but the *T. cristinae* DNA sequence data from  
260 Gompert et al. (2022) have been archived on NCBI's SRA (PRJNA356801) and the pheno-  
261 typic data and genotype estimates for *T. cristinae* are available on Dryad ([https://doi.](https://doi.org/10.5061/dryad.m905qfv26)  
262 [org/10.5061/dryad.m905qfv26](https://doi.org/10.5061/dryad.m905qfv26)).

## 263 **Code availability**

264 Computer code for the new simulation analysis is available from github ([https://github.](https://github.com/zgompert/TimemaPolygenicSelection/blob/main/SimMsLD.R)  
265 [com/zgompert/TimemaPolygenicSelection/blob/main/SimMsLD.R](https://github.com/zgompert/TimemaPolygenicSelection/blob/main/SimMsLD.R)).

## 266 **Author contributions**

267 ZG, JF and PN wrote and revised the manuscript.

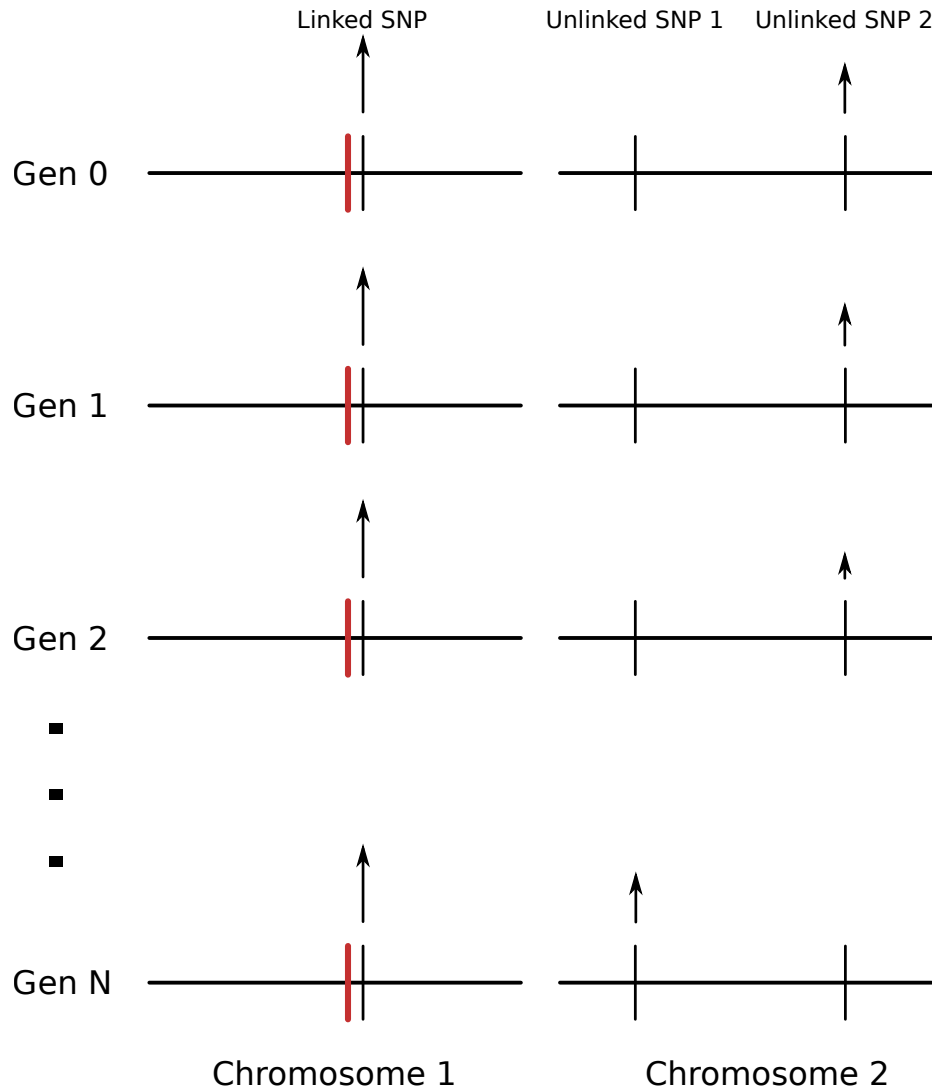


Figure 1: Hypothetical illustration of short-term, long-distance indirect selection. The vertical red line denotes a genetic variant causally affecting fitness (i.e., under direct selection). Vertical black lines denote a linked SNP and two unlinked SNPs. Arrows denote the degree of linkage disequilibrium (LD) between each SNP and the causal genetic variant, with larger arrows denoting higher levels of LD. Because of limited recombination, the linked SNP remains in high LD with the causal variant for an extended period of time (i.e., up to generation N, where N might denote tens or hundreds of generations). In contrast, because of free recombination, LD between unlinked SNP 2 and the causal variant decays over several generations. Nonetheless, during these few generations, indirect selection causes some of the allele frequency change at this locus to be deterministic and directional. Moreover, in this hypothetical example, LD is later created between the causal variant and a different unlinked SNP (unlinked SNP 1) because of some combination of drift, gene flow and selection. Thus, LD with unlinked variants is ephemeral but also constantly recreated, leading to perpetual indirect selection.

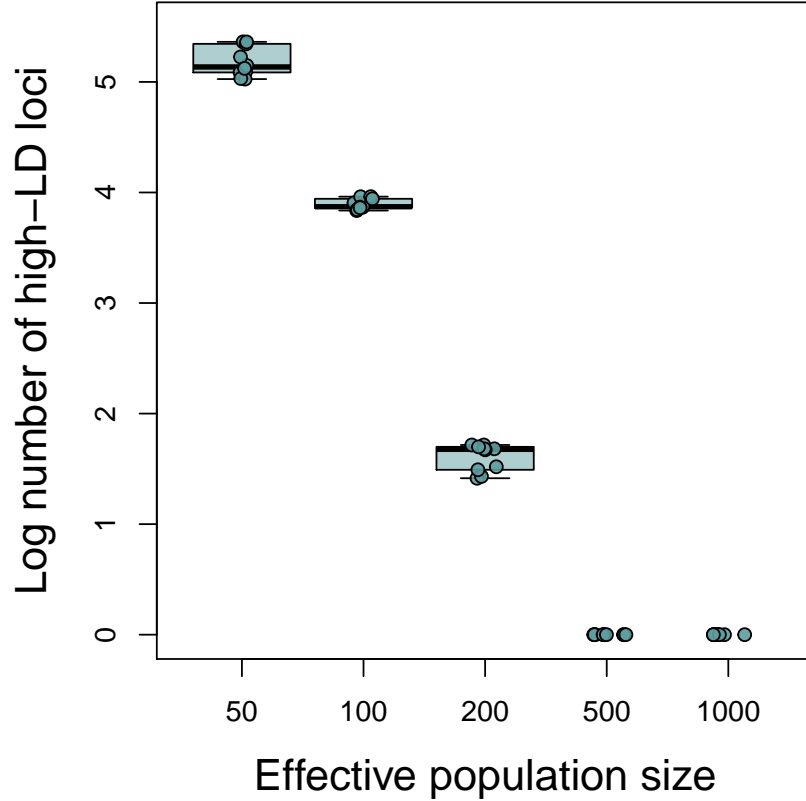


Figure 2: Boxplots with overlain points show the number of genetic loci (e.g., SNPs) in high linkage disequilibrium (LD), here defined as  $r^2 > 0.1$ , with a focal locus in simulations of genetic drift. Results are shown for each of 10 simulations with effective population sizes of 50, 100, 200, 500 or 1000 individuals. Numbers are shown on a  $\log_{10}$  scale, with the exception that all simulations with effective sizes of 500 and 1000 resulted in 0 high-LD SNPs ( $\log_{10}(0)$  is  $-\infty$  but 0 values are shown instead). Simulations involved a focal locus with minor allele frequency of 0.32 (our estimate of the minor allele frequency for *Mel-Stripe* in the *T. cristinae* population) and 7 million unlinked SNPs with allele frequencies drawn from the estimated allele frequencies at genome-wide SNPs in *T. cristinae*. We used binomial sampling to draw genotypes at each locus for each individual with independent draws across loci (i.e., we simulated unlinked loci). LD was then computed as the squared genotypic correlation ( $r^2$ ). We conducted these simulations in R (version 4.0.2).