Experimental introgression in *Drosophila*: asymmetric postzygotic isolation associated with chromosomal inversions and an incompatibility locus on the X chromosome

Running title: Experimental introgression in *Drosophila*

Authors:
Poikela N.¹, Laetsch D. R.², Kankare M.¹, Hoikkala A. *¹*, & Lohse K.*²

¹ Department of Biological and Environmental Science, University of Jyväskylä, Finland
² Institute of Evolutionary Biology, University of Edinburgh, United Kingdom

Corresponding author:
Noora Poikela, Department of Biological and Environmental Science, P.O. Box 35, FI-40014 University of Jyväskylä, Finland
E-mail: noora.p.poikela@gmail.com
Phone: +358 40 5383527

* Shared last authorship

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Abstract

Interspecific gene flow (introgression) is an important source of new genetic variation, but selection against it can reinforce reproductive barriers between interbreeding species. We used an experimental approach to trace the role of chromosomal inversions and incompatibility genes in preventing introgression between two partly sympatric Drosophila virilis group species, D. flavomontana and D. montana. We backcrossed F₁ hybrid females from a cross between D. flavomontana female and D. montana male with the males of the parental species for two generations and sequenced pools of parental strains and their reciprocal 2nd generation backcross (BC₂mon and BC₂fla) females. Contrasting the observed amount of introgression (mean hybrid index, HI) in BC₂ female pools along the genome to simulations under different scenarios allowed us to identify chromosomal regions of restricted and increased introgression. We find no deviation from the HI expected under a neutral null model for any chromosome for the BC₂mon pool, suggesting no evidence for genetic incompatibilities in backcrosses towards D. montana. In contrast, the BC₂fla pool showed high variation in the observed HI between different chromosomes, and massive reduction of introgression on the X chromosome (large X-effect). We find that this observation is compatible with reduced recombination combined with at least one dominant incompatibility locus residing within the X inversion(s). Overall, our study suggests that genetic incompatibilities arising in chromosomal inversions can play an important role in speciation.

Keywords: chromosomal inversions, experimental evolution, genetic incompatibilities, hybridization, introgression, X-effect
**Introduction**

Interspecific gene flow (introgression) is an important source of genetic variation for adaptation to new environments (Abbott et al., 2013; Anderson & Hubricht, 1938; Lewontin & Birch, 1966). At the same time, selection against introgression at certain loci acts to maintain barrier loci and protect species’ integrity from the negative effects of hybridization (Barton & Bengtsson, 1986; Ravinet et al., 2017; Servedio & Noor, 2003; Wu, 2001). The patterns of genomic divergence and the permeability of species boundaries in certain genomic regions provides valuable insights into the genomic regions that contribute to speciation (Harrison & Larson, 2014). However, we still lack a good understanding of how barrier genes are arrayed within the genome, how effectively and in what generation they restrict introgression, and what kind of role chromosomal inversions and sex chromosomes play in maintaining genetic barriers (Butlin, 2005; Coughlan & Matute, 2020; Coyne & Orr, 2004; Faria & Navarro, 2010; Gompert, Lucas, Nice, & Buerkle, 2012; Nosil & Feder, 2012).

Speciation in isolation (allopatriy), occurring via drift or indirect effects of selection, can lead to the “incidental” establishment of intrinsic genetic incompatibilities (Coyne & Orr, 2004; Tang & Presgraves, 2009). These incompatibilities generally involve negative epistatic interactions between two or more loci, where new alleles arising in one or both of the interacting lineages function well in their own genetic background, but interact negatively with the alleles of other species in hybrids (Bateson-Dobzhansky-Muller incompatibilities, BDMIs or DMIs; Coyne & Orr, 2004; Orr, 1995; Presgraves, 2010b). Lack of gene flow may also increase the fixation probability of meiotic drive loci (loci that manipulate meiotic process to favour their own transmission) and their suppressor loci within each population and drive the genomic divergence of these populations (Crespi & Nosil, 2013). Compared to allopatric speciation, where both BDMIs and neutral differences between species are expected to build up randomly along the genome, divergence with gene flow leads to clusters of species- or population-specific loci that are sheltered from recombination (Abbott et al., 2013; Butlin, 2005; Felsenstein, 1981). Accordingly, an accumulation of BDMIs between species may be drastically different with and without gene flow. Importantly, in the presence of gene flow BDMIs can only accumulate if they are favoured by selection (Bank, Bürger, & Hermisson, 2012).

Chromosomal inversions are a major factor rearranging the genome and inducing changes in gene interactions and expression patterns (Hoffmann & Rieseberg, 2008; Kirkpatrick & Barton, 2006; Sturtevant, 1921; Dobzhansky, 1940). Inversions may gain a fitness advantage and spread through conspecific populations if they reduce recombination between co-adapted genes (Kirkpatrick & Barton, 2006; Navarro & Barton, 2003). Once inversions have become fixed between species, they can generate postzygotic isolation in several ways. They can prevent interspecific gene flow directly by inducing problems in chromosome pairing during meiosis, which can lead to malformed gametes and reduced hybrid fertility and viability (Coyne & Orr, 2004; Hoffmann & Rieseberg, 2008; Rieseberg, 2001). However, these problems are partially avoided in *Drosophila*, since malformed gametes remain in the polar nuclei and do not enter the developing gametes (Hoffmann & Rieseberg, 2008; Sturtevant & Beadle, 1936). Perhaps more importantly, the limited recombination across inverted regions, particularly near inversion breakpoints and within overlapping inversions, facilitates the build-up of BDMIs via divergent selection and/or drift (Fishman, Stathos, Beardsley, Williams, & Hill, 2013; Khadem, Camacho, & Nóbrega, 2011; Mcgau & Noor, 2012; Navarro & Barton, 2003; Noor, Grams, Bertucci, & Reiland, 2001). Thus, species-specific inversions harbouring BDMIs may act as strong barriers to gene flow (Hoffmann & Rieseberg, 2008; Noor et al., 2001).

The disproportionate involvement of sex chromosomes in reproductive isolation in many systems is captured by two general observations: Haldane’s rule – the increased F1 inviability and sterility of the heterogametic sex compared to the homogametic sex (Haldane, 1922; Orr, 1997; Turelli & Orr, 2000) – and the large X-effect – the fact that the X chromosome shows a disproportionalately large effect on the sterility and inviability of backcross hybrids (Masly & Presgraves, 2007; Turelli & Orr, 2000). Explanation
for both observations often presume recessivity of X-linked alleles, which can lead to more pronounced effects in hemizygous than in heterozygous hybrids ("Dominance theory"; Coyne & Orr, 2004; Turelli & Orr, 1995, 2000) and/or rapid evolution of X-linked alleles facilitating BDMIs as a byproduct ("Faster X evolution"; Charlesworth, Campos, & Jackson, 2018; Charlesworth, Coyne, & Barton, 1987). The X chromosome has also been suggested to be enriched for genes that create postzygotic isolation in hybrids compared to autosomes (Coyne, 2018). In particular, meiotic drive loci are more frequent on the X than on autosomes, and incompatibilities between drivers and their suppressors in hybrids may generate problems in hybrid development (Courret, Chang, Wei, Montchamp-Moreau, & Larracuente, 2019; Crespi & Nosil, 2013; Crown, Miller, Sekelsky, & Hawley, 2018).

Pairwise BDMIs may involve substitutions in both diverging lineages, or derived substitutions in one lineage and preserved ancestral alleles in another lineage (Barbash, Awadalla, & Tarone, 2004; Cattani & Presgraves, 2009; Coyne & Orr, 2004). BDMIs can also result from cumulative effects of many small incompatibilities or from a single incompatibility between two complementary genes, and the complexity of the incompatibility interaction does not reflect the severity of the barrier (Orr, 1995; Presgraves, 2010a). Importantly, and in contrast to interactions within a locus where a dominant allele masks a recessive allele, in epistatic interactions between different loci a dominant allele at one locus may interact with dominant or recessive alleles at other loci. Epistatic interactions involving dominant alleles are of special interest in the context of BDMIs but have received less attention than BDMIs involving recessive alleles.

Two closely-related species of the Drosophila virilis group, D. montana and D. flavomontana, provide an excellent test case for studying the evolution of BDMIs. The species originate from the Rocky Mountains of North America, where the divergence of the montana complex species (D. flavomontana, D. lacicola and D. borealis) most likely occurred (Hoikkala & Poikela, 2022; Patterson, 1952; Throckmorton, 1982). D. montana has expanded around the northern hemisphere, whereas D. flavomontana has remained in North America (Hoikkala & Poikela, 2022). D. montana lives generally in colder environments and uses different host trees than D. flavomontana (Patterson, 1952; Throckmorton, 1982). Reproductive barriers between D. montana females and D. flavomontana males are nearly complete, while in the reciprocal cross strong postzygotic isolation is accompanied by prezygotic barriers of variable strength (Poikela et al., 2019). Regardless of these barriers, the two species can be crossed to obtain backcross progenies in both parental directions (Poikela et al., 2019), and interspecific hybrids have reportedly been found in nature (Patterson, 1952; Throckmorton, 1982). Our recent demographic modelling shows that the species have diverged ~3 Mya, with low levels of postdivergence gene flow from D. montana to D. flavomontana (Poikela et al., in prep.). Moreover, we found in these species several alternatively fixed inversions, which were already present in their common ancestor, and which may have contributed to the build-up and maintenance of adaptive traits and reproductive barriers by restricting gene flow between the evolving lineages (Poikela et al., in prep.).

The goal of this study was to determine which genomic regions are likely to accommodate dominant BDMIs in hybrids between D. montana and D. flavomontana, paying special attention to fixed inversions and the X chromosome. We investigated BDMIs between these species experimentally by sequencing pools of D. montana females from an allopatric population and D. flavomontana females from a (presently) parapatric population, as well as pools of 2nd backcross generation (BC2) females in both directions (Fig. 1). We identified chromosomal regions with decreased and increased introgression by quantifying the amount of introgressed genetic material (mean hybrid index, HI) along the genome in both backcross pools. We then compared the observed HI to the distribution of chromosome-wide HI in in silico replicates of this “introgress-and-resequence” experiment under contrasting assumptions about the presence and location of BDMIs. Since this experimental design involved only backcross females, only BDMIs involving a dominant allele could affect allele frequencies in the pool while recessive-recessive BDMIs were masked (Table 1). Our main questions were:
2 Materials and methods

2.1 Fly material

We collected fertilised *D. montana* females from Seward, Alaska, USA (60°09’N; 149°27’W) and *D. flavomontana* females from Livingston, Montana, USA (45°20’N; 110°36’W) in 2013. The distance between the sites is ~3000 km. Alaskan *D. montana* can be regarded as an allopatric population, as *D. flavomontana* has not been found above 54°N (Poikela et al., 2019). In contrast, the *D. flavomontana* population from Montana can be regarded as a parapatric, as the two species are known to coexist in the Rocky Mountains, even though we found only *D. flavomontana* on the collecting site (Poikela et al., 2019).

Progenies of wild-caught *D. montana* and *D. flavomontana* females were used to establish isofemale strains, which were maintained in continuous light and 19 °C for about 23 generations (~3 years) in the University of Jyväskylä (Finland) prior to their use in the present study. We performed the crossing experiment using flies of *D. montana* strain SE13F37 and *D. flavomontana* strain MT13F11. For the crosses, the flies were sexed under light CO₂ anaesthesia within three days after emergence, when they were still virgins. Males and females were transferred into fresh malt-vials once a week and used in the crossing experiments at age 20 ± 2 days when they were sexually mature (Salminen & Hoikkala, 2013).

2.2 Crossing experiment

We started the crossing experiment by performing a single-pair cross between *D. flavomontana* female and *D. montana* male, as reciprocal cross is not successful. Our crossing design (outlined in Fig. 1) only involved hybrid females because F₁ males are largely sterile (Pääällysaho, Aspi, Liimatainen, & Hoikkala, 2003; Poikela et al., 2019), and because *Drosophila* males lack recombination (crossing-over) in meiosis.

The initial cross produced seven F₁ females, which were backcrossed towards both parental species: four were mated to *D. montana* males and three to *D. flavomontana* males. The 1st backcross generation females (BC₁mon and BC₁fla females) were backcrossed to the same paternal species as in the previous generation to obtain BC₂mon and BC₂fla females (82 females in both directions). BC₂ females were collected within three days after their emergence and stored in -20 °C for DNA extractions.

2.3 Fertility of BC₁ females

We defined the fertility of BC₁ females (BC₁mon and BC₁fla females) by checking whether they produced progeny after mating with a *D. montana* or *D. flavomontana* male (Fig. 1). BC₁ females that produced no progeny were considered sterile. We used a one-sample Student’s t-test (t-test function) to test whether the BC₁ females from the reciprocal crosses showed reduced fertility, when the expected fertility is 1. We also compared the fertility of BC₁ females between the reciprocal crosses to define possible asymmetries (BC₁mon vs. BC₁fla), using a generalised linear model (GLM) with Binomial distribution (1=fertile, 0=sterile) (glm function). All analyses were conducted in base R v1.2.1335-1 and R studio v3.6.1.

2.4 Pool-sequencing, mapping, and variant calling

We made DNA extractions from four pools, one pool of each parental strain (*D. montana* SE13F37 and *D. flavomontana* MT13F11) and pools for the two 2nd generation backcrosses (BC₂mon and BC₂fla). Each
pool consisted of 82 females. We used cetyltrimethylammonium bromide (CTAB) solution with RNAse treatment, Phenol-Chloroform-Isosamyl alcohol (25:24:1) and Chloroform-Isosamyl alcohol (24:1) washing steps and ethanol precipitation. Nextera library preparation and 150 bp Illumina paired-end sequencing were performed at Edinburgh Genomics, UK. Illumina paired-end reads of all four samples were quality-checked with FastQC v0.11.8 (Andrews 2010) and trimmed for adapter contamination and low-quality bases using fastp v0.20.0 (Chen, Zhou, Chen, & Gu, 2018). After filtering, the total number of reads per pool varied from 152 to 174 million, and the mean length and insert size peak being 141-143bp and 150bp, respectively (Table S1). The mean coverage of the pools varied from 170 to 220 (Table S1).

All analyses were based on reads mapped to a D. montana chromosome-level genome assembly with full gene annotation (Poikela et al. in prep.). Filtered Illumina reads of each sample were mapped to the unmasked reference genome using BWA mem (Burrows-Wheeler Aligner) v0.7.17 with read group information (Li & Durbin, 2009). The alignments were sorted with SAMtools v1.10 (Li et al., 2009) and PCR duplicates marked with sambamba v0.7.0 (Tarasov, Vilella, Cuppen, Nijman, & Prins, 2015). The obtained BAM-files were used for variant calling with the softmasked version of the reference genome using freebayes parallel v1.3.1-dirty (Garrison & Marth, 2012) with --no-population-priors --hwe-priors-off --use-mapping-quality --ploidy 2 --theta 0.02 --haplotype-length -1.

Variant calling detected a total of 8,876,483 variants. To normalise the representation of variants, the resulting VCF-file was processed with vt normalize (Tan, Abecasis, & Kang, 2015). The variants were filtered for quality and SNPs (single nucleotide polymorphism) using bcftools filter and view v1.9 (Li, 2011). We chose only biallelic SNPs with a minimum depth of 80 to reliably calculate allele frequencies and to minimize potential reference bias. These quality filtering steps resulted in a total of 5,047,746 SNPs.

2.5 Inversion breakpoints

The breakpoints of alternatively fixed inversions of D. montana and D. flavomontana on the X chromosome and chromosomes 2L, 4 and 5 were obtained from Poikela et al. (in prep.). The presence of the inversions in Illumina samples of parental pools was verified by passing the respective BAM-files to Delly v0.8.1 (Rausch et al., 2012), which identifies structural variants based on paired-end read orientation and split-read evidence. The inversion breakpoints were also confirmed visually by checking the orientation and insert size around each breakpoint in the Interactive Genomics Viewer (Thorvaldsdóttir, Robinson, & Mesirov, 2012) (Example plot shown in Fig. S1). Inversion breakpoints are shown in Fig. 3, Fig. 4 and Table S2.

2.6 Genetic differentiation, hybrid index and the types of genetic incompatibilities

The expected amount of genetic material transferred from one species into the other (hybrid index, HI) halves with every backcross generation (Fig. 1). Thus, in the pool of 2nd backcross generation hybrid females, the genome-wide HI is expected to be 12.5% in the absence of BDMIs (Fig. 1). However, given the random inheritance of chromatids in gametes and the randomness of cross-over locations, we expect substantial variation around the expected mean HI, even in the absence of BDMIs.
Figure 1. Illustration of the crossing experiment showing the inheritance of sex chromosomes (inheritance of autosomes is similar to that of female X chromosomes). F\textsubscript{1} females, produced in a single-pair cross between D. flavomontana (fla) female and D. montana (mon) male, were backcrossed to either D. flavomontana or D. montana male. In the next generation, each BC\textsubscript{1} female was mated with a male of its paternal species. In every generation, the expected amount of genetic material that is transferred from the gene pool of one species into the gene pool of another one (introgression) is halved (red percentages). Thus, under a null neutral model, we expect a mean HI of 12.5% for the BC\textsubscript{2} pools that were sequenced. Note that recombination occurring in the gametes produced by F\textsubscript{1} and BC\textsubscript{1} females creates variation in the expected amount of HI. For simplicity, the figure shows products of only one cross-over event that has occurred in each backcross direction.

To estimate the amount of introgression in the BC\textsubscript{2} pools, we computed the HI in both pools along the genome based on species-diagnostic SNPs (variants that are differentially fixed between the parental pools). Allele frequencies for each SNP in all four pools were calculated by dividing “Alternate allele observation count (AO)” by “the total read depth (DP)” (relative to D. montana reference genome). Diagnostic variants were defined as SNPs with allele frequency 1 in one parental pool and 0 in the other one (1 = all reads supporting the alternate allele, 0 = all reads supporting the reference allele). The total number of SNPs that were differentially fixed between the parental species was 1,109,701.

We compared collinear and inverted parts of each chromosome in terms of the density of diagnostic SNPs. Each chromosome was divided into 200kb non-overlapping windows (55-147 windows per chromosome depending on the chromosome length), and the number of diagnostic SNPs in each window was counted using a custom script [https://github.com/vihoikka/SNP_mapper/blob/main/snp_binner.py](https://github.com/vihoikka/SNP_mapper/blob/main/snp_binner.py). The data was analysed using a generalised linear model (glm function) with a Poisson distribution, where the number of window-wise SNPs was used as a response variable, and different chromosomes and different chromosomal partitions (collinear, inverted) as explanatory variables. The analyses were performed in base R using R v1.2.1335-1 and R studio v3.6.1.

Using the diagnostic SNPs, we calculated the mean hybrid index (HI) separately for different chromosomes for BC\textsubscript{2}fla and BC\textsubscript{2}mon pools. We also estimated the fraction of sequence without any introgressed material (HI = 0%) separately for each chromosome for both pools. Finally, we plotted HI in non-overlapping windows of 400 SNPs for each chromosome and BC\textsubscript{2} pool using a custom script [https://github.com/vihoikka/SNP_mapper/blob/main/datasmoother.py](https://github.com/vihoikka/SNP_mapper/blob/main/datasmoother.py). In principle, recombination breakpoints involving the two ancestral backgrounds (Fisher junctions; Fisher, 1954) should be visible...
as step changes in the HI of each pool. Assuming on average one cross-over per chromosome and female meiosis, the expected number of recombination breakpoints per chromosome generated during the experiment is given by the total number of females (nBC1 + nBC2; Table S3) contributing to each pool (96 and 104 for BC1,mon and BC1,fla pools, respectively). Note that the number of junctions between *D. montana* and *D. flavomontana* ancestral material is lower since not all cross-over events in BC1 females are junctions. In practice, however, the resolution especially for breakpoints that are unique to a single BC1 individual (which correspond to a change in allele frequency of 1/82) is limited by the randomness in sequencing coverage of the pool.

Given that this experiment was started with a single-pair cross between the parental species and continued with repeated backcrosses between hybrid females and parental males, all backcross individuals inherited a maximum of one allele per locus from the donor species (Fig. 1). Thus, the genomes of BC individuals are a mosaic of two types of tracts: i) homozygous for the genetic background of the recipient species or ii) heterozygous between species. This limits the types of BDMIs that can be expressed (Table 1). Dominant-dominant pairwise BDMIs arise already in the F1 generation and, if severe, can cause sterility/inviability in both sexes. Recessive-recessive pairwise BDMIs are not possible in our experiment even if they were X-linked since i) all BC individuals involved in the experiment were females (no hemizygosity), and ii) the expression of these incompatibilities would require homozygous tracts for both species (Fig. 1). Hence, dominant-recessive BDMIs are the only strong postzygotic barriers that we expect to detect in this study.

Table 1. BDM model for incompatibilities (see Coyne & Orr, 2004). Here gene A1 of one species interacts negatively with gene B1 of another species. Underscore represents any allele, and it does not change the outcome. Note that dominance refers to an allele’s effect on fitness on a hybrid genetic background, and it does not necessarily assume dominance of alleles on their normal background within species.

| dominant-dominant incompatibility (both loci act dominantly): | A1_ B2_ hybrids are affected in the F2 generation |
| recessive-recessive incompatibility (both loci act recessively): | A1A2B1B2 hybrids are affected in the F2 generation |
| dominant-recessive incompatibility (A1 acts dominantly, B2 recessively): | A1_ B1B2 hybrids are affected in backcross generations |

### 2.7 Simulating the backcross and re-sequence experiment

Given the stochastic nature of inheritance of chromatids in gametes and the randomness of cross-over locations in meiosis, we expected substantial variation in the HI around the expectation of 12.5% (Fig. 1). To evaluate whether the observed mean HI of each chromosome deviates significantly from that expected under simple models of introgression with or without inversions and/or extreme BDMIs, we simulated the crossing experiment under three different scenarios using Mathematica (Wolfram Research, Inc., version 11.02 Champaign, IL). All simulations were conditioned on the number of BC2 females each BC1 female contributes to the pool (Table S3). We also assumed one cross-over per female per chromosome in meiosis (a map length of 50cM). Given that the experiment involves two generations of crosses between hybrid females and pure parental males, our simulation only tracks the haplotype of female gametes contributing to BC1 and BC2 individuals. All *in silico* backcross experiments were simulated, separately for each chromosome, 10,000 times to obtain 5% and 95% quantiles for the mean HI.

First, we simulated the experiment under a simple null model of neutral introgression, i.e. assuming no BDMIs and no cross-over suppression due to inversions (SIM1, Fig. 2A). Second, we simulated the
experiment similarly under neutrality, but including the breakpoint locations of inversions that are
alternately fixed between \textit{D. montana} and \textit{D. flavomontana}. This was done simply by disallowing cross-
over events within inverted regions (inversions breakpoints in Table S2), i.e. we did not attempt to
include interchromosomal effects (SIM2, Fig. 2B). Third, we simulated the experiment under a model
that assumes a single BDMI at a random position within the inverted part of the chromosome (SIM3,
Fig. 2C). This single locus cannot be introgressed beyond the \textit{F} \textsubscript{1} generation, i.e. BC\textsubscript{1} and BC\textsubscript{2} females that
are heterozygous for this locus are not produced. Note that while we refer to this as a BDMI for
simplicity, we did not explicitly simulate pairwise incompatibilities. Thus, this locus can be regarded as
a BDMI involving a dominant allele on the introgressing background (donor species) that is incompatible
with one or more recessive alleles in the recipient background.

![Figure 2](image)

**Figure 2.** Introgression experiment was simulated under different scenarios. Example plots of simulated hybrid
indices (HI) (A) under neutrality, (B) in the presence of neutral inversions, and (C) in the presence of inversions
with a single dominant BDMI. For simplicity, here simulations were run 10 times.

### 3 Results

#### 3.1 BC\textsubscript{1} females from the backcrosses towards \textit{D. flavomontana} showed stronger genetic
incompatibilities / postzygotic isolation than the ones from the backcrosses towards \textit{D. montana}

The proportion of fertile females in the BC\textsubscript{1} generation was 75\% and 42\% for the BC\textsubscript{1}mon and BC\textsubscript{1}fla
hybrids, respectively, and significantly reduced in both reciprocal crosses when comparing to the
expected fertility of 1 (BC\textsubscript{1}mon: \textit{t}\textsubscript{19} = -2.52, \textit{P} = 0.021; BC\textsubscript{1}fla: \textit{t}\textsubscript{54} = -8.67, \textit{P} = 8.371e \textsuperscript{-12}). Furthermore, the
proportion of fertile BC\textsubscript{1}mon females (75\%) was significantly higher than that of BC\textsubscript{1}fla females (42\%)
(GLM, \textit{z}_{1,73} = -2.45, \textit{P} = 0.015; Fig. S2). These findings show that while both crosses suffer from BDMLs
affecting female fertility, these incompatibilities are more pronounced in backcrosses towards \textit{D.
flavomontana} than towards \textit{D. montana} (asymmetric postzygotic isolation, or unidirectional
incompatibilities in the sense of Turelli & Moyle, 2007).

#### 3.2 Genetic divergence between \textit{D. montana} and \textit{D. flavomontana} has accumulated within inverted
chromosome regions especially on the X chromosome

The density of SNPs (SNPs divided into equal-sized bins) that were differentially fixed between \textit{D.
montana} and \textit{D. flavomontana} parental pools was higher on the X chromosome than on any of the
autosomes (\textit{P} < 0.001; Fig. 3; Table S4). For each chromosome containing inversions, the density of fixed
differences was higher in inverted than in collinear regions of the genome (\textit{P} < 0.001; Fig. 3; Table S5),
as expected due to the reduction in recombination within inverted regions (note that chromosomes 2R
and 3 have no inversions).
The mean amount of introgression (hybrid index, HI) of hybrids backcrossed to *D. montana* (BC$_{mon}$) did not deviate significantly from the neutral expectation of 12.5% for any chromosome (SIM1) (Fig. 4A; Fig. 5). Moreover, the number of SNPs that showed no introgression was low, 0.03-0.22%, across the entire genome (Table S6).

In contrast, BC$_{fla}$ hybrids showed a significant deviation in mean HI from the neutral scenario (SIM1) for all chromosomes, except chromosome arms 2L and 2R (Fig. 4; Fig. 5B; Table S6). The mean HI was significantly decreased on the X and 4$^{th}$ chromosome and significantly increased on the 3$^{rd}$ and 5$^{th}$ chromosome compared to the neutral expectation (SIM1) (Fig. 4B; Fig. 5; Table S6). Interestingly, the reduced introgression on the 4$^{th}$ chromosome and the increased introgression on the 5$^{th}$ chromosome could be explained by the reduction in cross-over due to inversions present on these chromosomes, without invoking any selection acting on incompatibilities (SIM2) (Fig. 4, 5E, 5F). Under this scenario, the mean HI showed no deviation from the expectation of 12.5% under neutrality but had an increased variance across simulation replicates (Fig. 5E, 5F). Also, the number of SNPs that showed no introgression varied from 0.12% to 1.22% for chromosomes 2L, 2R, 4 and 5 (Table S6).

The observed decrease in the mean HI of BC$_{fla}$ hybrids on the X chromosome could not be explained solely by a reduction in cross-over rate due to inversions (Table S6, Fig. 4, 5C). Instead, our simulations show that the drastic reduction in mean HI on the X chromosome is compatible with a single dominant incompatibility locus residing within the X inversions (SIM3) (Fig. 4, 5G). In other words, the data are consistent with a dominant X chromosomal *D. montana* allele that interacts negatively with autosomal homozygous recessive *D. flavomontana* alleles. Intriguingly, 30 % of the differentially fixed SNPs between the species on the X chromosome showed no introgression, emphasising the strength of the X-effect (Table S6).

Finally, we conclude that the increase in mean HI observed in BC$_{fla}$ hybrids on the 3$^{rd}$ chromosome cannot be explained by any of the simple scenarios we simulated. Given that we either assumed neutrality or a single dominant incompatibility locus, which is maximally deleterious in the BC$_{1}$ and BC$_{2}$ females, this is not surprising. The number of SNPs showing no introgression was only 0.18% for the 3$^{rd}$ chromosome.
Figure 4. Observed hybrid index (HI) of 2nd backcross generation female pools towards D. montana (BC2mon) and D. flavomontana (BC2fla) in windows of 400 non-overlapping SNPs along the genome. For chromosome 2 the left (2L) and right (2R) arms are separated by a metacentric centromere. The black horizontal line represents the expected amount of introgression, HI = 12.5 %, under neutrality. Vertical lines represent species-specific D. flavomontana (yellow) and D. montana (blue) chromosomal inversions. Solid and dashed vertical lines show breakpoints of different inversions.

Figure 5. Hierarchical representation of the most meaningful simulations (10,000 replicates/simulation) of the 2nd generation backcross experiments towards D. montana (BC2mon) and D. flavomontana (BC2fla). The grey area of each figure represents Bonferroni corrected 5% and 95% quantiles and the space between them (regions beyond the area are statistically significant). Simulations under neutrality (SIM1) and the observed mean hybrid index (HI) of each chromosome for (A) BC2mon pool and (B) BC2fla pool. Simulations under neutral inversions (SIM2) and observed mean HI of BC2fla pool for (C) the X chromosome, (D) chromosome 3, (E) chromosome 4, and (F)
4 Discussion

4.1 Postzygotic barriers between D. montana and D. flavomontana show asymmetry in their strength

We have previously shown that pre- and postzygotic barriers between D. montana females and D. flavomontana males are practically complete, while both types of barriers between D. flavomontana females and D. montana males are weaker (Poikela et al., 2019). In crosses between D. flavomontana females and D. montana males, F₁ hybrid males are sterile, but roughly half of the F₁ females are fertile (Poikela et al., 2019). Accordingly, here we backcrossed fertile F₁ females with males of both parental species, and observed a clear asymmetry in the strength of postzygotic barriers between the two backcross directions. BC₁ hybrid females born from the backcrosses between F₁ females and D. montana males showed rather high fertility, and the genetic incompatibilities observed in BC₂ females had no detectable effect. In contrast, when backcrossing F₁ hybrid females with D. flavomontana males, more half of the BC₁ females were sterile, and BC₂ females showed signs of strong BDMIs. This asymmetry could be a consequence of a history of unidirectional introgression from D. montana into D. flavomontana in nature (Poikela et al., in prep.), if it had induced selection against introgression at certain loci especially within the X chromosomal inversions, but homogenised genetic divergence on collinear regions. This kind of pattern in the permeability of species boundaries have been found to contribute to speciation also in other species (Harrison & Larson, 2014).

It is surprising that introgression has not occurred from D. flavomontana to D. montana in nature, given that backcrossing towards D. montana (BC₁ mon) was relatively successful in this study. The most obvious reason for this discrepancy is that laboratory experiments may not reveal all reproductive barriers relevant in wild populations. For example, hybrids may have problems in mate choice in the wild, or they may face challenges to feed or reproduce on species-specific host trees. Moreover, also the male hybrids regain fertility in backcross generations (data not shown), which may contribute to introgression in nature. Finally, in this study we used a D. montana population that is allopatric to D. flavomontana, while BDMIs may well be stronger between D. montana and D. flavomontana populations living in close contact.

4.2 The role of inversions and the X chromosome in reducing recombination and introgression from D. montana to D. flavomontana (BC₂ fla pool)

Inversions have suggested to contribute to speciation, if three criteria are met: closely related species must carry alternatively fixed inversions, the inversions must suppress recombination, and this suppression of recombination facilitates reproductive isolation (Faria & Navarro, 2010). We have recently identified several alternatively fixed inversions in D. montana and D. flavomontana, and shown that these inversions have increased genetic divergence and lower historical introgression compared to collinear chromosome regions (Poikela et al., in prep.). In the present study, we show that these inversions have increased number of alternatively fixed SNPs compared to collinear regions, which is in agreement with their increased genetic divergence shown in Poikela et al. (in prep.). We also show that inversions effectively suppress recombination in hybrid individuals across a large swathe of the...
genome (Fig. 4). Finally, we find that the drastic reduction in introgression on the X chromosome can be explained by inversions that are associated with at least one dominant X chromosomal D. montana incompatibility allele interacting negatively with recessive autosomal D. flavomontana alleles. This negative epistatic interaction could cause the observed low hybrid fertility, and supports the idea that inversions act as strong barriers to gene flow by facilitating the establishment of BDMIs (Hoffmann & Rieseberg, 2008; Navarro & Barton, 2003; Noor et al., 2001).

While the involvement of the X chromosome in hybrid problems may not be surprising (see e.g. Masly & Presgraves, 2007; Tao, Chen, Hartl, & Laurie, 2003), the fact that it involves a dominant incompatibility locus is. The “dominance theory” (e.g. Turelli & Orr, 1995, 2000), which aims to explain the disproportionate role of the X chromosome in hybrid incompatibilities, relies on the presence of recessive incompatibilities on the X and therefore cannot explain our result. However, the “dominance theory”, as well as the “faster-male theory” and dosage compensation (reviewed in Coyne, 2018; Presgraves, 2008), can still explain the hybrid male sterility previously observed in crosses between D. flavomontana and D. montana (Poikela et al., 2019). Accumulation of meiotic drive elements on the X chromosome could be another plausible explanation for the large X-effect in general (reviewed in Patten, 2018), but is unlikely in our system for two reasons. First, meiotic drivers should increase their own transmission in both backcross directions, leading to decreased introgression in the BC2fla pool and increased introgression in the BC4mon pool, which we did not see. Second, meiotic drive systems described in Drosophila are typically involved in sperm killing and not in female sterility (Courret et al., 2019). Although cytoplasmic incompatibilities have been detected in other montana complex species of the Drosophila virilis group (Patterson, 1952; Throckmorton, 1982), they are not likely to play a major role in these crosses since all hybrids had D. flavomontana cytoplasm (and crosses were more unsuccessful in this direction). Finally, the large effect of the X we find could potentially be explained by “faster X evolution”, based on the idea that selection increases the frequency of advantageous recessive alleles more effectively on the X chromosome than on autosomes, irrespectively of whether the incompatibilities themselves are recessive (Charlesworth et al. 1987, 2018). Also, the X chromosome could simply contain more genes that are prone to create postzygotic isolation than those on the autosomes (Coyne, 2018).

Several autosomes showed deviations from the expected hybrid indices in the BC2fla pool in the present study. Based on our simulations, the reduced and increased introgression on the 4th and 5th chromosomes, respectively, could be explained by inversions’ ability to restrict recombination which increases the variance in chromosome-wide HI. However, if we calculate the expected allele frequencies for a dominant–recessive BDMI by hand for the first two backcross generations, the allele frequencies (i.e. HI) after selection would be 1/22 (4.5%) for the dominant and 2/11 (18.2%) for the recessive D. montana allele in the BC2fla pool (see Fig. S2). These frequencies are close to the observed frequencies on chromosomes 4 (5.4%) and 2L (19.8%), respectively. It is therefore tempting to speculate that a pairwise BDMI loci exist on these chromosomes. Finally, the 3rd chromosome, which lacks species-specific inversions, showed a drastic increase in introgression, and was not explained by any of our simulations. We note that our simulations did not consider an interchromosomal effect, where inversions may trigger an increase in recombination on other freely recombining chromosomes (Crown et al., 2018; Stevison, Hoehn, & Noor, 2011). However, this would only decrease the variance in HI on chromosomes lacking fixed inversions and so cannot explain the increase in HI for chromosome 3 in the BC2fla pool.

In future research, combing the crosses with quantitative trait loci (QTL) analyses might help to link BDMIs to e.g. specific genes (Johnson, 2010) or gene duplicates or transpositions (Bikard et al., 2009; Masly, Jones, Noor, Locke, & Orr, 2006). BDMI genes could also be searched by tracing the expression-phenotypes in interspecific hybrids in these genes (Landry et al., 2005). However, recombination suppression of inversions presents a challenge for mapping BDMIs, and would potentially require a
complex reversion of the X chromosomal inversions with genome editing tools, and repeating the
current experiment to narrow down the regions of reduced introgression (Hopkins, Tyukmaeva,
Gompert, Feder, & Nosil, 2020). Overall, finding the exact loci driving species’ isolation may be difficult,
as BDMIs are often complex and co-evolve with rapidly evolving heterochromatic DNA (Satyaki et al.,
2014).

5 Conclusions

“Introgress-and-resequence” studies that combine interspecific backcrosses with genome-wide
analyses and simulations are an effective approach for identifying BDMIs, in particular those involving
dominant alleles. Our study supports the idea that inversions aid the accumulation of BDMIs due to
reduced recombination, and shows that strong BDMIs coupled with suppressed recombination
effectively restrict introgression beyond the inverted part of the genome in the first two backcross
generations. We conclude that the large X-effect we observed in our experiment may result from at
least one dominant incompatibility locus residing within several overlapping inversions. If the design
were extended to study interspecific F2 hybrids, assuming that the F1 female and male hybrids are viable
and fertile, one could investigate recessive-recessive BDMIs in the same way. Overall, we provide a novel
framework for investigating the role of inversions and the X chromosome as genetic barriers to
introgression, which we hope will encourage similar studies on a larger number of species and strains.

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Data Accessibility

Raw reads will be made publicly available in the SRA (BioProject XXX), and phenotypic data and
Mathematica notebook including simulations in Dryad at the time of publication.

Author Contributions

KL, AH and NP designed the study. NP performed the hybrid backcrosses and analysed the genomic data
with input from KL and DRL. KL performed the simulations. AH and MK supervised and funded the
research. NP, AH and KL drafted the manuscript and all authors finalised it.

Conflict of interest

The authors declare no conflict of interest.

ORCID

Noora Poikela: 0000-0002-4627-9647
Dominik R. Laetsch: 0000-0001-7887-0186
Maaria Kankare: 0000-0003-1541-9050
Anneli Hoikkala: 0000-0001-5407-7992
Konrad Lohse: 0000-0001-9918-058X

Ethics declaration

Neither species is endangered, and the flies were collected along watersides on public lands outside
National and State parks, where insect collecting does not require permits in the USA (The Wilderness
Act of 1964, section 6302.15).
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