Locally Adaptive Inversions Modulate Genetic Variation at Different Geographic Scales in a Seaweed Fly

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Abstract

Across a species range, multiple sources of environmental heterogeneity, at both small and large scales, create complex landscapes of selection, which may challenge adaptation, particularly when gene flow is high. One key to multidimensional adaptation may reside in the heterogeneity of recombination along the genome. Structural variants, like chromosomal inversions, reduce recombination, increasing linkage disequilibrium among loci at a potentially massive scale. In this study, we examined how chromosomal inversions shape genetic variation across a species range and ask how their contribution to adaptation in the face of gene flow varies across geographic scales. We sampled the seaweed fly Coelopa frigida along a bioclimatic gradient stretching across 10° of latitude, a salinity gradient, and a range of heterogeneous, patchy habitats. We generated a chromosome-level genome assembly to analyze 1,446 low-coverage whole genomes collected along those gradients. We found several large nonrecombining genomic regions, including putative inversions. In contrast to the collinear regions, inversions and low-recombining regions differentiated populations more strongly, either along an ecogeographic cline or at a fine-grained scale. These genomic regions were associated with environmental factors and adaptive phenotypes, albeit with contrasting patterns. Altogether, our results highlight the importance of recombination in shaping adaptation to environmental heterogeneity at local and large scales.

Key words: structural variants, population genomics, local adaptation, diptera, environmental associations.

Introduction

Across its range, a species experiences variable environmental conditions at both small and large geographic scales. This environmental heterogeneity makes local adaptation a complex process driven by multiple dimensions of selection and constrained by the distribution of genetic diversity within the genome and the intensity of gene flow acting on it (Savolainen et al. 2013; Tigano and Friesen 2016). Recombination plays a complex role in mediating this process (Stapley et al. 2017). On the one hand, recombination reduces Hill-Robertson interference, allowing natural selection to act on single loci (Otto and Barton 2001; Roze and Barton 2006). On the other hand, recombination homogenizes populations and reshuffles coadapted or locally adapted groups of alleles (Charlesworth and Charlesworth 1979; Lenormand and Otto 2000). Hence, the landscape of recombination influences adaptive trajectories, depending on the distribution of environmental heterogeneity, epistasis,

(Charlesworth and Charlesworth 1979; Lenormand and Otto 2000; Yeaman 2013), and is expected to modulate the geographic distribution of adaptive and nonadaptive genetic diversity (Ortiz-Barrientos and James 2017; Stevison and McGaugh 2020).

Chromosomal inversions are major modifiers of the recombination landscape, whereby recombination between the standard and inverted arrangements is reduced in heterokaryotypes (Sturtevant 1917; Hoffmann et al. 2004). A single species can have multiple polymorphic inversions, each of them covering hundreds of kilobases or megabases, thus their impact can be widespread across the genome (Wellenreuther and Bernatchez 2018). For instance, five polymorphic inversions are present worldwide in Drosophila melanogaster (Kapun and Flatt 2019) and maize (Zea mays) harbors a 100 Mb inversion (Fang et al. 2012). The last decade has shown that such inversion polymorphisms occur in a wide range of species and has brought important insights into the

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adaptive role of inversions (Hoffmann and Rieseberg 2008; Wellenreuther and Bernatchez 2018; Mérot, Oomen, et al. 2020). Inversions with a large effect on complex multitrait phenotypes, such as life-history, behavior, and color patterns, confirm that arrangements can behave as haplotypes of a "supergene," linking together combinations of alleles within each arrangement (Joron et al. 2011; Schwander et al. 2014; Kirubakaran et al. 2016; Wellenreuther and Bernatchez 2018; Yan et al. 2020). Inversions are also notable for their associations with segregation distorters, involving epistatic selection which favors linkage between coadapted alleles at interacting loci (Sturtevant and Dobzhansky 1936; Fuller et al. 2020). Likewise, covariation between inversion frequencies and environmental variables, whether spatial, temporal, or experimental (Dobzhansky 1948; Schaeffer 2008; Kapun et al. 2016; Kirubakaran et al. 2016; Faria et al. 2019; Kapun and Flatt 2019; Huang and Rieseberg 2020) is consistent with selection for the suppression of recombination between locally adaptive loci (Kirkpatrick and Barton 2006; Charlesworth and Barton 2018) and/or coadaptive epistatic interactions between loci (Dobzhansky and Dobzhansky 1970; Charlesworth and Charlesworth 1973). Hence, when investigating adaptation with respect to multiple scales and at multiple sources of environmental variation, it is important to examine the role of large inversions or any recombination suppressors.

Coelopa frigida is a seaweed fly that inhabits piles of rotting seaweed, so-called wrackbeds (fig. 1), on the east coast of North America and in Europe. Coelopa frigida is known to harbor one large inversion on chromosome I (hereafter called Cf-Inv(1)) that is polymorphic in Europe and America (Butlin,

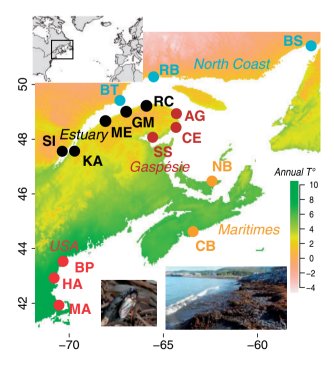


Fig. 1. Coelopa frigida sampling across an environmental gradient. Map of the 16 sampling sites, colored by geographic region. The background of the map displays the gradient of annual mean air temperature. The insert shows the location of the study area at a wider scale. Photos show *C. frigida* and its habitat of seaweed beds.

Collins, et al. 1982; Mérot et al. 2018), as well as four additional large polymorphic inversions described in one British population (Aziz 1975). The inversion Cf-Inv(1) encapsulates 10% of the genome and has two arrangements: α and β . These alternative Cf-Inv(1) arrangements have opposing effects on body size, fertility, and development time, a combination of traits that results in different fitnesses depending on the local characteristics of the wrackbed (Butlin, Read, et al. 1982; Day et al. 1983; Butlin and Day 1985; Edward and Gilburn 2013; Wellenreuther et al. 2017; Berdan et al. 2018; Mérot, Llaurens, et al. 2020). Almost nothing is known about the other inversions but, given that a large fraction of the C. frigida genome is affected by polymorphic inversions, one can expect that these inversions play a significant role in structuring genetic variation and contribute to local adaptation. Spatial genetic structure and connectivity in C. frigida remain poorly described, although occasional long-distance migration bursts have been documented and regular dispersal is expected between nearby subpopulations occupying discrete patches of wrackbed (Egglishaw 1960; Dobson 1974). Coelopa frigida occupies a wide climatic range of temperature (temperate to subarctic zones) as well as salinity (from freshwater to fully saline sites). Furthermore, C. frigida experiences high variability in the quality and the composition of its wrackbed habitat (Egglishaw 1960; Dobson 1974). These sources of habitat heterogeneity vary at both large and local geographic scales, for which, depending on the scale of dispersal, a linked genomic architecture may be favorable.

In the present study, we investigated how chromosomal inversions contribute to local adaptation across different scales of environmental heterogeneity, and how they shape genetic diversity. Using the seaweed fly C. frigida as a biological model, we adopted a systematic approach for localizing multiple chromosomal inversions and analyzed genetic variation across several dimensions of environmental variation including a 1,500 km climatic gradient, a salinity gradient, and fine scale, patchy habitat variation (fig. 1). We built the first reference genome assembly for C. frigida and sequenced 1,446 whole genomes at low coverage. Using this comprehensive data set, we analyzed patterns of genetic polymorphism along the genome to identify putative inversions. As connectivity between populations of C. frigida was previously unknown, we examined its geographic structure with respect to single nucleotide polymorphism (SNP) markers. Finally, we tested genotype-environment and genotype-phenotype associations to determine the genomic architecture of adaptation to various sources of environmental variation acting at different geographic scales.

Results

To facilitate our analyses, we built the first reference genome assembly for *C. frigida* using a combination of long-read sequencing (PacBio) and linked-reads from 10x Genomics technology. A high-density linkage map (28,639 markers segregating across six linkage groups [LGs]) allowed us to anchor and orientate more than 81% of the genome into five large chromosomes (LG1 to LG5) and one small sex

chromosome (LG6). This karyotype was consistent with previous cytogenetic work on *C. frigida* (Aziz 1975) and with the six Muller elements (A = LG4, B = LG5, C = LG2, D = LG3, E = LG1, F = LG6, supplementary fig. S1, Supplementary Material online), which are usually conserved in Diptera (Vicoso and Bachtrog 2015; Schaeffer 2018). The final assembly included six chromosomes and 1,832 unanchored scaffolds with an N50 of 37.7 Mb for a total genome size of 239.7 Mb. This reference had a high level of completeness, with 96% (metazoan) and 92% (arthropods) of universal single-copy orthologous genes completely assembled. It was annotated with a highly complete transcriptome (87% complete BUSCOs in the arthropods) based on RNA-sequencing of several ontogenetic stages and including 35,999 transcripts.

To analyze genomic variation at the population-scale, we used low coverage (\sim 1.4X) whole-genome sequencing of 1,446 flies from 16 locations along the North American Atlantic coast (88–94 adult flies/location). Sampled locations spanned a North-South gradient of 1,500 km, over 10° of latitude, a pronounced salinity gradient in the St Lawrence Estuary, and a range of habitats with variable seaweed composition and wrackbed characteristics (fig. 1 and supplementary table S1, Supplementary Material online). After alignment of the 1,446 sequenced individuals to the reference genome, we analyzed genetic variation within a probabilistic framework accounting for low coverage (ANGSD, Korneliussen et al. 2014) and reported 2.83 million SNPs with minor allelic frequencies higher than 5% for differentiation analyses.

Two Large Chromosomal Inversions Structure Intraspecific Genetic Variation

Decomposing SNPs genotype likelihoods through a principal component analysis (PCA) revealed that the 1st and 2nd principal components (PCs) contained a large fraction of genetic variance, respectively, 21.6% and 3.9%, and allowed us to display the 1,446 flies as nine discrete groups (fig 2A). Along PC1, the three groups corresponded to three genotypes of the inversion Cf-Inv(1) ($\alpha\alpha$, $\alpha\beta$, and $\beta\beta$), as identified using two diagnostic SNPs (Mérot et al. 2018) with, respectively, 100% and 98.3% concordance (supplementary table S2, Supplementary Material online). Along PC2, three distinct groups were identified that corresponded neither to sex nor geographic origins, and thus possibly represented three genotypes for another polymorphic inversion.

To assess which regions of the genome reflected the patterns observed in the whole-genome PCA, we performed local PCA on windows of 100 SNPs along each chromosome and evaluated the correlation between PC1 scores of each local PCA and PCs scores of the global PCA (fig. 2C). PC1 was highly correlated with a region of 25.1 Mb on LG 1, indicating the genomic position of the large Cf-Inv(1) inversion (table 1). PC2 was highly correlated with a smaller region of 6.9 Mb on LG4 (fig. 2D), consistent with the hypothesis of an inversion, hereafter called Cf-Inv(4.1). Several other characteristics were consistent with the hypothesis that these two regions are inversions. First, inside these regions, linkage disequilibrium (LD) was very high when considering all individuals, but low

within each group of homokaryotypes (fig. 2B). This indicates that recombination is limited between the arrangements but occurs freely in homokaryotypes bearing the same arrangement. Second, F_{ST} was very high between homokaryotes in the inverted region (Cf-Inv(1) $\alpha\alpha$ vs. $\beta\beta$: 0.75, Cf-Inv(4.1) AA vs. BB: 0.51, fig. 2C) compared with low values in the rest of the genome (Cf-Inv(1) $\alpha\alpha$ vs. $\beta\beta$: 0.002, Cf-Inv(4.1) AA vs. BB: 0.001, fig. 2D). Third, the intermediate group on the PCA was characterized by a higher proportion of observed heterozygotes for SNPs in the inverted region than the extreme groups, confirming that this is probably the heterokaryotypic group (supplementary fig. S2, Supplementary Material online).

Nucleotide diversity, as measured by π , was similar between karyotypic groups along the genome, and higher in the heterokaryotypes than in the homokaryotypes in inverted regions (fig. 2C and D). For both inversions, nucleotide diversity was comparable between homokaryotes. Absolute nucleotide divergence between arrangements was strong in inverted regions (table 1 and supplementary fig. S3, Supplementary Material online). Assuming a mutation rate comparable to Drosophila (5 \times 10⁻⁹ mutations per base per generation [Assaf et al. 2017]), and approximately five to ten generations per year, we thus estimated, from absolute divergence at noncoding regions that the arrangements split at least 180,000 to 376,000 years ago for Cf-Inv(1) and at least 61,000 to 134,000 years ago for Cf-Inv(4.1).

Coelopa frigida Exhibit Other Regions Including Nonrecombining Haplotypic Blocks

To further examine the heterogeneity of genetic structure along the genome, we reanalyzed the local PCAs using a method based on multidimensional scaling (MDS) that identifies clusters of PCA windows displaying a common pattern. This method has been previously used to identify and locate nonrecombining haplotypic blocks (Li and Ralph 2019; Huang et al. 2020; Todesco et al. 2020). Besides the aforementioned Cf-Inv(1) and Cf-Inv(4.1) inversions, which caused the 1st and 2nd axis of the MDS, we identified five outlier genomic regions across the different MDS axes (fig. 3 and supplementary fig. S4, Supplementary Material online). In all five regions, a large proportion of variance was captured along the 1st PC (>50%), and LD was high (fig. 3A).

Two regions on LG4 represented convincing putative inversions of 2.7 and 1.4 Mb, respectively. In both regions, the PCA displayed three groups of individuals with high clustering confidence; the central group contained a high proportion of heterozygotes and the extreme groups were differentiated (fig. 3E and supplementary fig. S5, Supplementary Material online). Within these two regions, nucleotide diversity was comparable between haplogroups and the absolute divergence (d_{XY}) between homokaryotypes was lower than for Cf-Inv(1) and Cf-Inv(4.1), suggesting younger inversions that could have diverged as recently as 6,000 to 68,000 years ago. Karyotype assignment was the same between the two putative inversions, indicating that they are either tightly linked or belong to a single inversion. Two lines of evidence support the hypothesis that these are two

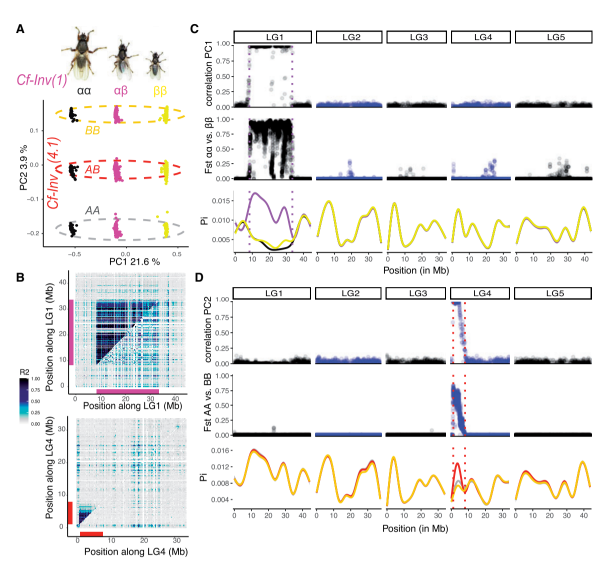


Fig. 2. Two large chromosomal inversions structure within species genetic variability. (A) PCA of whole-genome variation. Individuals are colored by karyotypes at the inversion Cf-Inv(1), as determined previously with an SNP marker (Mérot et al. 2018). Ellipses indicate secondary grouping along PC2. (B) LD in LG1 and LG4. The upper triangles include all individuals and the lower triangles include homokaryotes for the most common arrangement for each inversion. Bars represent the position of the inversions. The color scale shows the 2nd higher percentile of the R^2 value between SNPs summarized by windows of 250 kb (C) Along the genome, correlation between PC1 scores of local PCAs performed on windows of 100 SNPs and PC1 scores of the PCA performed on the whole genome; F_{ST} differentiation between the two homokaryotypes of Cf-Inv(1) in sliding-windows of 25 kb; and nucleotide diversity (π) within the three karyotypic groups of Cf-Inv(1) smoothed for visualization. Dashed lines represent the inferred boundaries of the inversion Cf-Inv(1) (D) Correlation between the two homokaryotypes of Cf-Inv(4.1) in sliding windows of 25 kb; and nucleotide diversity (π) within the three karyotypic groups of Cf-Inv(4.1) smoothed for visualization. Dashed lines represent the inferred boundaries of the inversion Cf-Inv(4.1).

inversions. First, the high density of linkage map markers and the non-null recombination rate across this area of 50 cM provided confidence in the genome assembly and supported a gap of 5 Mb between the two inversions. Moreover, previous cytogenetic work showed that one chromosome of *C. frigida* exhibits a polymorphic inversion on one arm (possibly *Cf-Inv*(4.1)) and, on the other arm, two polymorphic inversions, which rarely recombine (Aziz 1975). Both inversions were subsequently analyzed together and called *Cf-Inv*(4.2) and *Cf-Inv*(4.3).

The other three regions, spanning 6.8 Mb on LG2, 6.3 Mb on LG3, and 16.7 Mb on LG5, represented complex areas that

behaved differently from the rest of the genome. Recombination was locally reduced, both in the linkage map and in wild populations, as indicated by strong LD (fig. 3A and B). These three regions were all highly heterogeneous; within each region, nucleotide diversity showed highly contrasting pattern across subregions (fig. 3C). A fraction of these subregions exhibited low nucleotide diversity, which may correspond to centromeric or pericentromeric regions (fig. 3C and supplementary fig. S6, Supplementary Material online), as well as a high density of transposable elements, such as LINEs or LTRs (supplementary fig. S7, Supplementary Material online). However, these low diversity subregions were

Table 1. Name, Position, and Characteristics of the Putative Inversions and Regions Appearing as Cluster of Outlier Windows in the Local PCA Analysis.

Name	Status	Chr.	Start	Stop	Size (MB)	d_{XY}		
Cf-Inv(1)	Known inversion	LG1	8,342,182	33,487,673	25.1	1.84% [1.80-1.88]		
Cf-Inv(4.1)	Probable inversion	LG4	1,088,816	7,995,568	6.9	0.64% [0.61-0.67]		
Cf-Inv(4.2)	Probably two linked inversions	LG4	22,421,881	25,145,365	2.7	0.079% [0.061-0.096]		
Cf-Inv(4.3)	·	LG4	30,622,035	31,991,919	1.4	0.32% [0.31-0.34]		
Cf-Lrr(2)	Low-recombination region	LG2	14,083,320	20,869,940	6.8			
Cf-Lrr(3)	Low-recombination region	LG3	7,486,933	13,829,649	6.3			
Cf-Lrr(5)	Low-recombination region	LG5	15,940,464	32,665,323	16.7			

Note.—For putative inversions, absolute nucleotide divergence (d_{XY}) in noncoding regions was calculated between homokaryotypic groups and corrected by the mean of nucleotide diversity (π) within homokaryotypic groups by windows of 25 kb. Numbers between square brackets indicate confidence intervals drawn by bootstrapping windows of 25 kb.

interspersed with subregions of high diversity, particularly on LG5 (fig. 3C). Some of those high diversity subregions also corresponded to clusters of outlier windows in the local PCA analysis and appeared as nonrecombining haplotypic blocks of medium size (1–2 Mb) in partial LD (supplementary figs. S8–S10, Supplementary Material online). In the absence of more information about the mechanisms behind the reduction in recombination, we consider those three regions of the genome to be simply "low recombining regions" (subsequently called *Cf-Lrr(2)*, *Cf-Lrr(3)*, *Cf-Lrr(5)*). Accordingly, the fraction of the genome subsequently called "collinear" excluded both these regions and the inversions (*Cf-Inv(1)*, *Cf-Inv(4.1)*, *Cf-Inv(4.2)*, and *Cf-Inv(4.3)*).

Geographic Structure Shows Distinctive Signals in Inverted and Low-Recombining Regions

Geography also played a major role in structuring genetic variation. Our 3rd PC, which explained 1.4% of variance, represented genetic variation along the North-South gradient (fig. 4A). Differentiation between pairs of populations, measured as F_{ST} on a subset of LD-pruned SNPs, also followed the North-South gradient but was globally weak ($F_{ST} = 0.003$ to 0.016, supplementary fig. S11, Supplementary Material online). We also detected a strong signal of isolation by distance (IBD) when examining the correlation between genetic distances and Euclidean distances among the 16 populations (R^2 = 0.45, F = 97, P < 0.001, supplementary table S3, Supplementary Material online). Considering least cost distances along the shorelines instead of Euclidian distances between locations improved the model fit ($R^2 = 0.63$, F = 199, P < 0.001, $\Delta AIC = 47$, table 2 and supplementary table S3, Supplementary Material online). This supports a pattern of isolation by resistance (IBR, see Materials and Methods), in which dispersal occurs primarily along the coastline and is limited across the mainland or the sea.

These IBD and IBR patterns varied significantly along the genome. When considering all SNPs, pairwise differentiation was more heterogeneous ($F_{\rm ST}=0.002$ to 0.021, fig. 4B) and IBR was much weaker, albeit significant ($R^2=0.19,\,F=29,\,P<0.001$) than when considering LD-pruned SNPs or collinear SNPs. We thus calculated pairwise $F_{\rm ST}$ between pairs of populations based on different subsets of SNPs, either from each inversion, from each low-recombining region, or from the collinear genome.

All of the inversions exhibited increased differentiation between populations in comparison with the collinear genome (supplementary table S3 and fig. S12, Supplementary Material online). However, the global geographic patterns differed between inversions. In the inverted region Cf-Inv(1), there was no association between genetic and geographic distances (fig. 4B and table 2), a result that significantly contrasts with the collinear genome (supplementary fig. \$13, Supplementary Material online). This result was due to highly variable pairwise genetic differentiation between populations in the inverted region Cf-Inv(1). Conversely, genetic differentiation between geographic populations in the inverted regions of LG4 showed significant IBD/IBR patterns with a significantly steeper slope of regression between genetic and geographic distances compared with collinear regions (fig. 4B and C, table 2 and supplementary table S4 and fig. S13, Supplementary Material online). The divergence between northern and southern populations was mirrored by a sharp and significant latitudinal cline of inversion frequencies, ranging from 0.27 to 0.75 for Cf-Inv(4.1) (GLM: z = -8.1, P < 0.001, $R^2 = 0.41$) and from 0.02 to 0.26 for Cf-Inv(4.2/4.3) (GLM: z = -6.6, P < 0.001, $R^2 = 0.37$). The association between latitude and inversion frequency was significantly stronger than for randomly chosen SNPs with similar average frequencies (fig 4D and supplementary figs. S14 and S15, Supplementary Material online).

Although the entire genome (with the exception of inversion *Cf-Inv(1)*) showed IBD and IBR, it was significantly increased in two of the three low-recombining regions compared with the collinear regions. When compared with collinear regions of the same size, the slope of the regression between genetic and geographic distances was significantly steeper for *Cf-Lrr(2)* and *Cf-Lrr(5)* but not for *Cf-Lrr(3)* (fig. 4C, table 2 and supplementary table S4 and fig. S13, Supplementary Material online). Overall, the geographic differentiation in the four inverted regions and two low-recombining regions showed patterns differing from the collinear genome, indicating the influence of processes other than the migration-drift balance, possibly at different geographic scales for *Cf-Inv(1)* vs. others.

Adaptive Diversity Colocalizes with Inversions and Low-Recombining Regions

To investigate putative patterns of adaptive variation in *C. frigida*, we analyzed the association between SNP frequencies

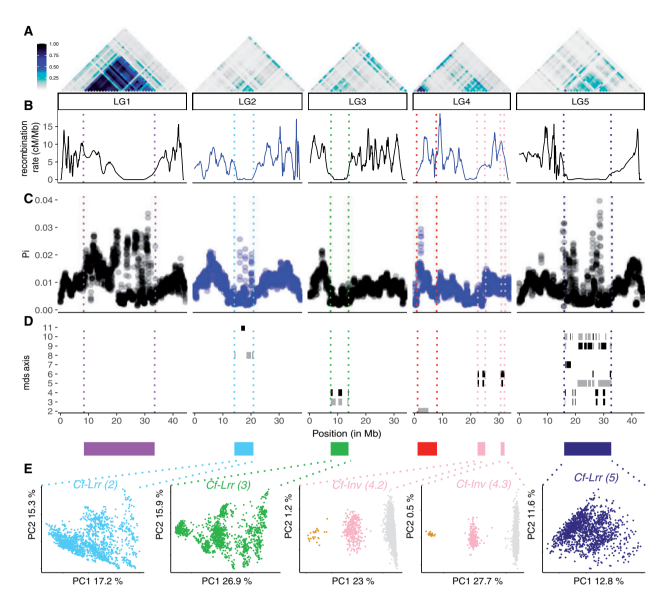


Fig. 3. Detecting other regions exhibiting non recombining haplotypic blocks. (A) LD across the five major chromosomes expressed as the 2nd higher percentile of the R^2 value between SNPs summarized by windows of 1 Mb. (B) Recombination rate (in cM/Mb) inferred from the linkage map, smoothened with a loess function accounting for 10% of the markers. (C) Nucleotide diversity (π) by sliding windows of 100 kb (step 20 kb) averaged across the different geographic populations. (D) Position along the genome of clusters of local PCA windows scored as outliers (>4 SD) along each axis of the MDS, at the upper end in black, and the lower end in gray. Colored rectangles indicate the position of the inversions and the regions of interest gathering outlier clusters or putative inversions. Dashed lines represent their inferred boundaries across all plots. (E) PCA performed on SNPs located in each region of interest. For the two regions on LG4 that appear as two linked putative inversions (*Cf-Inv*(4.2) and *Cf-Inv*(4.3)), three clusters were identified with high confidence and colored as putative homokaryotes and heterokaryotes. The same colors are used in both regions since karyotyping was consistent across all individuals.

and environmental variables at large (thermal latitudinal gradient and salinity gradient in the St. Lawrence R. Estuary) and local (abiotic and biotic characteristics of the wrackbed habitat) spatial scales (fig. 1 and supplementary fig. S16 and table S1, Supplementary Material online). Analyses with two different genotype-environment association (GEA) methods (latent factor mixed models and Bayesian models) showed consistent results, highlighting high peaks of environmental associations and large clusters of outlier SNPs in the inverted or low-recombining regions (fig. 5A–E, table 3 and supplementary table S5 and figs. S17 and S18, Supplementary Material online). However, different inversions were

implicated depending on environmental factor and spatial scale. We considered SNPs consistently identified as outliers across both analyses to be putatively adaptive.

At a large geographic scale, associations with climatic variation along the latitudinal gradient showed a strong excess of outlier SNPs in the four inversions and the low-recombining regions of LG2 and LG5. These regions exhibited two to five times more outliers than expected by chance (table 3) with particularly strong peaks of environmental association (Bayes factor [BF] >50, fig. 5A), and a signal significantly stronger than for random blocks of collinear genome of the same size (supplementary fig. S19, Supplementary Material online).

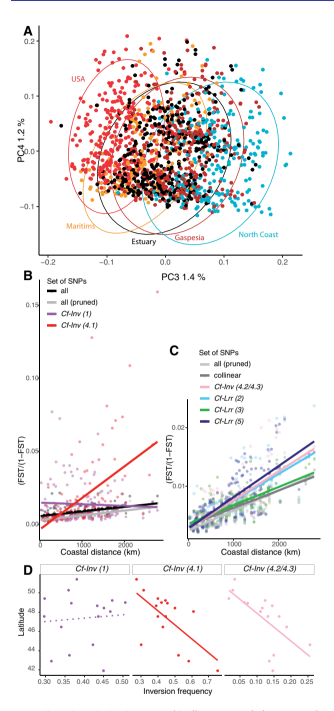


Fig. 4. Genetic variation is geographically structured along a North-South gradient and displays IBR. (A) Third and 4th PCs of a PCA on whole-genome variation. Individuals are colored by their geographic region, as in fig. 1. (*B* and *C*) IBR displayed as the association between genetic distance ($F_{ST}/(1-F_{ST})$) and the distance by the least-cost path following the coast. Colors denote the subset of SNPs used for the calculation of the F_{ST} . The results are displayed in two panels with different y scales to better display the lower values. (*D*) Latitudinal variation of inversion frequencies.

However, this was not the case for *Cf-Lrr*(3). These results were consistent whether or not the model was controlled by the geographic population structure (supplementary figs. S17 and S18, Supplementary Material online). Variation of frequencies of *Cf-Inv*(4.1) and *Cf-In*(4.2/4.3) was also

significantly associated with climatic variation, when considered as single loci (GLM: Cf-Inv(4.1): z=-7.76, P<0.001; Cf-Inv(4.2/4.3): z=-6.45, P<0.001, with the model explaining 36% and 37% of the variance in inversion frequency, supplementary fig. S20, Supplementary Material online). Variation along the salinity gradient, which also spanned variation in tidal amplitude, was significantly associated with a more limited number of SNPs. A large excess of such outliers were found in Cf-Lrr(3) and Cf-Lrr(5) (table 3), the only two regions in which the signal of association was stronger than in the collinear genome (supplementary fig. S19, Supplementary Material online).

At a finer geographic scale, outlier SNPs associated with wrackbed abiotic characteristics (depth, temperature, and salinity) were strongly enriched in the inverted region Cf-Inv(1) with an odds ratio of 5, including outliers with very strong support (BF >20, fig. 4C). SNP associations with wrackbed abiotic characteristics were stronger than in collinear regions in Cf-Inv(1), and more marginal in Cf-Inv(4.2/4.3) (supplementary fig. S19, Supplementary Material online). This was mirrored by the Cf-Inv(1) frequency, which significantly covaried with wrackbed (GLM, z = 3.5, P < 0.001, $R^2 = 0.26$, supplementary fig. S20, Supplementary Material online). Variation in algal composition of the wrackbed, driven by the relative abundance of two dominant families of seaweed, Fucaceae or Laminariaceae, was significantly associated with widespread SNPs, although the inversion Cf-Inv(1) was overrepresented by an odds ratio of 1.4. Variation in secondary components of the substrate was more difficult to interpret as they covaried with latitude and temperature (supplementary fig. \$16, Supplementary Material online). Despite this, these secondary components were also associated with a large number of SNPs in Cf-Inv(1) and in Cf-Lrr(5) with odds ratio of 3.6 to 6 (fig. 5E), and a distribution of association scores significantly higher than in collinear blocks (supplementary fig. S19, Supplementary Material online).

Genotype-Phenotype Association

As wrackbed composition and Cf-Inv(1) are known to influence adult size (Butlin, Read, et al. 1982; Edward and Gilburn 2013), we used a genome-wide association study (GWAS) to uncover genetic variation associated with wing size. Among the 124,701 candidate SNPs identified by the GWAS, more than 99.8% were located in Cf-Inv(1) (fig. 5F). When variation in karyotype was removed (by running the analysis with only homokaryotypes), we found almost no candidate SNPs associated with size variation (0 for $\alpha\alpha$ individuals, and up to 3 SNPs when the FDR was lowered to P = 0.01 for the $\beta\beta$ individuals, supplementary fig. S21, Supplementary Material online). We ran gene ontology using two data sets: the candidates identified by GWAS and all genes present in Cf-Inv(1). Both analyses showed an enrichment in several biological processes all consistent with large differences in wing size and life history, such as morphogenesis, muscle development, or neural system development (supplementary tables S6 and S7, Supplementary Material online).

Given the extreme temperature range inhabited by *C. frigida* (temperate to subarctic), we also investigated thermal

Table 2. Association between Genetic Distance and Geographic Distances Measured as Least-Cost Distances along the Shoreline (IBR) for the Different Fractions of the Genome.

SNP subset	R ² adjusted	F	P value	Intercept	Slope coefficient	Comparison to collinear regions		
All	0.19	29.3	<0.001	0.0085	0.0020 [0.0013-0.0027]			
Collinear	0.54	138.6	< 0.001	0.0062	0.0019 [0.0015-0.0022]			
LD pruned	0.63	199.5	< 0.001	0.0057	0.0021 [0.0018-0.0024]			
Cf-Inv(1)	-0.01	0.3	0.59	0.0137	-0.0006 [-0.0032-0.0018]	_*		
Cf-Inv(4.1)	0.29	49.4	< 0.001	0.0172	0.0134 [0.0096-0.0172]	+*		
Cf-Inv(4.2/4.3)	0.50	121.5	< 0.001	0.0075	0.0030 [0.0025-0.0036]	+*		
Cf-Lrr(2)	0.44	95.4	< 0.001	0.0074	0.0028 [0.0023-0.0034]	+*		
Cf-Lrr(3)	0.49	113.1	< 0.001	0.0066	0.0019 [0.0016-0.0023]	n.s.		
Cf-Lrr(5)	0.55	147.2	< 0.001	0.0080	0.0033 [0.0028-0.0038]	+*		

NOTE.—Numbers between square brackets indicate the limits of the 95% distribution of the slope coefficient. The comparison to collinear regions displays the output of a full model comparing each region to the collinear genome, providing the direction and the significance (*) of the interaction term.

adaptation. We evaluated the recovery time after a chill coma in the F2s used to build the linkage map. Cold shock resistance localized to a quantitative trait locus (QTL) on LG4, which explained about 13% of the variation (supplementary fig. S22, Supplementary Material online). The main peak was located on LG4 around 25–28 Mb. This broad QTL encompassed multiple outliers SNPs associated with climatic variation, and multiple annotated genes, among them two heat shock proteins, which may represent relevant candidates for thermal adaptation (Uniprot P61604 at position 25,128,992 and P29844 at position 26,816,283). This peak was located between the two putative inversions *Cf-Inv*(4.2) and *Cf-Inv*(4.3), and there was a secondary peak at 8 MB, the putative breakpoint of *Cf-Inv*(4.1).

Discussion

Analyses of more than 1,400 whole genomes of C. frigida flies revealed four large chromosomal inversions affecting a large fraction of the genome (36.1 Mb, 15%), and three lowrecombining genomic regions. These megabases-long stretches of the genome appear to play a predominant role in shaping genetic variation across two large-scale environmental gradients as well as heterogeneous patchy habitats. Yet different inversions showed contrasting patterns, which may be related to different selective forces acting on them. In particular, the newly discovered inversions on LG4 displayed clinal variation along a geoclimatic gradient. In contrast, the largest inversion Cf-Inv(1) was associated with body size and covaried at a fine geographic scale with wrackbed habitat characteristics, confirming previous work (Day et al. 1983; Butlin and Day 1985; Mérot et al. 2018). Below, we discuss how our results provide new insights into the evolutionary role played by recombination-limited regions including inversions, and how our data suggest that those regions are involved in local adaptation at different geographic scales in the face of high gene flow.

Low-Coverage Sequencing Provides Insights into Genetic Variation across a Species Range and Individual Genomes

Studying all aspects of genetic variation across a species range is more accurate and powerful when sampling

encompasses both fine and coarse geographical scales across multiple environmental conditions. When searching for signatures of adaptation or putative inversions, high-density genetic markers are required to identify patterns (Fuentes-Pardo and Ruzzante 2017). This creates the need to balance effort across the number of samples, the portion of the genome sequenced (i.e., reduced representation or wholegenome sequencing), and the depth of sequencing. To maximize insights, we sequenced the whole genome of 1,446 wild-collected flies, but reduced individual coverage to about 1.4X. Simulations have shown that sequencing many samples at low depth (1X) provides robust estimates of population genetic statistics, namely allele frequencies, F_{ST} , and other population parameters, and may be more powerful than sequencing few samples at higher depth (Alex Buerkle and Gompert 2013; Lou et al. 2020). Consequently, this strategy has been used efficiently in a few pioneer studies in human genomics (Martin et al. 2021), conservation genomics (Therkildsen et al. 2019), population genomics (Clucas et al. Additionally, thanks to a low-cost barcoding library preparation (Therkildsen and Palumbi 2017), individual information was retained, which allowed parameters that require this information (LD, Hobs) to be accurately calculated as well as the use of phenotypic association studies. Importantly, allele frequencies were also unbiased by a priori or unbalanced pooling as may happen in pool-seq (Fuentes-Pardo and Ruzzante 2017), and any grouping could be subsequently chosen for the analyses.

Individual whole-genome sequencing at low coverage allowed us to uncover the genetic structure associated with inversions in *C. frigida* and to analyze environmental parameters and phenotypes potentially associated with those inversions. First, the large sample size brought power to make the most of a recently developed method of indirect inversion detection (Li and Ralph 2019; Huang et al. 2020). For instance, we would probably have missed the inversion(s) *Cf-Inv*(4.2/4.3) with smaller sample size, since the rare homokaryotype frequency was below 2% (26/1,446 individuals). Second, the high density of markers along the genome provided accurate locations for the major inversions although characterizing the exact breakpoints was

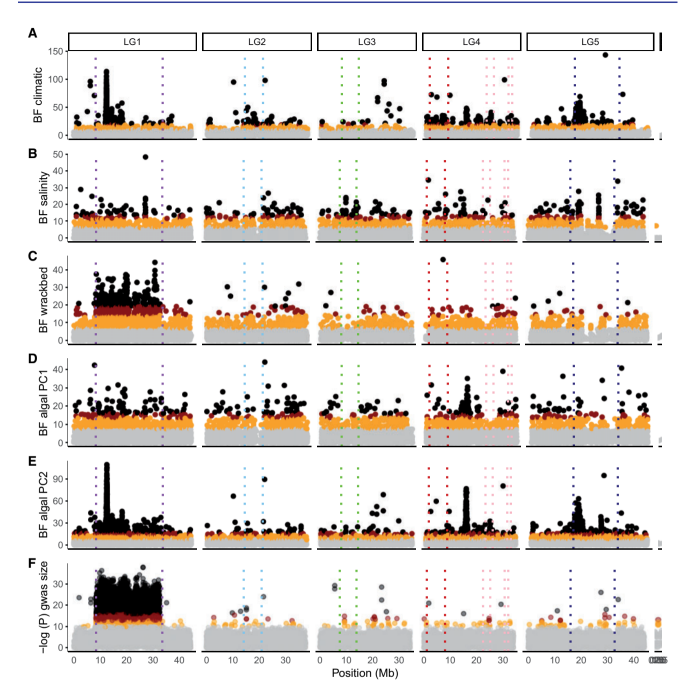


Fig. 5. Environmental and phenotypic associations. Candidate SNPs associated with (A) climatic variation along the North-South gradient, (B) salinity variation along the Estuarian gradient, (C) variations in abiotic characteristics of the wrackbed habitat, (D and E) variation in wrackbed algal composition. The Manhattan plot shows the Bayesian factor from the environmental association analysis performed in Baypass, controlling for population structure. (F) Candidate SNPs associated with wing size. The Manhattan plot shows the P values from the GWAS. Points are colored according to FDR (black: <0.00001, red: <0.0001, orange: <0.001). Dashed lines represent the inferred boundaries of inversions and low-recombining regions.

impossible without long-read sequencing (Ho et al. 2019). Third, the retention of individual information allowed us to split the data set into subgroups of karyotypes, as determined from the analyses of sequences, and to characterize LD, heterozygosity, nucleotide diversity, and the differentiation within and between karyotypes for all inversions. This aspect was critical to our study and allowed us to describe contrasting patterns from a geographic and ecological point of view.

Polymorphic Inversions Structure Within-Species Genetic Diversity

Despite the wide geographic area covered, the major factor shaping genetic variation in *C. frigida* was not geographic distance but chromosomal inversions. Along more than 1,500 km (or 3,000 km of coastal distance) between the most distant populations, geographic genetic differentiation was very weak (Maximal $F_{\rm ST} < 0.02$). This is much lower than other coastal specialized insects such as the saltmarsh beetle

Table 3. Genomic Repartition of Candidate SNPs Associated with Environmental Variables.

	Tested SNPs		Climate		Salinity		Bed abiotic characteristics		Algal composition (PC1)			Algal composition (PC2)					
	N	%	N	%	OR	N	%	OR	N	%	OR	N	%	OR	N	%	OR
All	1,155,978		3,635			509			780			372			2,740		
Collinear	814,279	70	556	15	0.2	301	59	0.8	163	21	0.3	254	68	1.0	390	14	0.2
Cf-Inv(1)	176,963	15	1474	41	2.6*	64	13	0.8	584	75	4.9*	77	21	1.4*	1494	55	3.6*
Cf-Inv(4.1)	57,323	5.0	480	13	2.7*	15	2.9	0.6	11	1.4	0.3	14	3.8	0.8	33	1.2	0.2
Cf-Inv(4.2/4.3)	17,019	1.5	111	3.1	2.1*	8	1.6	1.1	8	1.0	0.7	3	0.8	0.5	26	0.9	0.6
Cf-Lrr(2)	20,458	1.8	93	2.6	1.4*	6	1.2	0.7	9	1.2	0.7	3	0.8	0.5	15	0.5	0.3
Cf-Lrr(3)	16,313	1.4	11	0.3	0.2	28	5.5	3.9*	0	0.0	0.0	3	0.8	0.6	7	0.3	0.2
Cf-Lrr(5)	53,623	4.6	910	25	5.4*	87	17	3.7*	5	0.6	0.1	18	4.8	1.0	775	28	6.1*

NOTE.—Repartition of the candidate SNPs associated with each environmental variation using the combination of two GEA methods. *N* is the number of outliers SNPs located in a given region, % is the proportion of the outliers found in this region, and OR indicates the odds ratio. Values in bold with a star indicate significant excess of candidate SNPs in a Fisher exact test. Results obtained for each GEA method are presented in supplementary table S5, Supplementary Material online.

Pogonus chalceus ($F_{ST} \sim 0.2$, Van Belleghem et al. 2018) but comparable to small Dipterans with large distributions like D. melanogaster or Drosophila simulans, which typically exhibit F_{st} around 0.01-0.03, probably resulting from both high migration rate and large effective population size Ne (Machado et al. 2016; Kapun et al. 2020). Despite this weak genetic structure, we detected a strong signal of IBD indicating that dispersal among populations and subsequent gene flow decreases with distance. Furthermore, our analyses also showed that the least cost distance along the coastline better explained genetic variation than Euclidean distance. This IBR pattern probably results from a stepping stone dispersal process (Gandon and Rousset 1999) where the absence of suitable habitat patches in mainland and marine areas drives gene flow along the shore and constraints genetic connectivity.

In contrast with the overall weak geographic genetic structure, the frequencies of the different inversion arrangements were highly differentiated. Differentiation was restricted to the inverted regions, with fixed allelic differences between arrangements. Such a high genotypic divergence between alternative arrangements is comparable to many other ancient inversions (Hoffmann and Rieseberg 2008; Wellenreuther and Bernatchez 2018) and reflects the accumulation of neutral and non-neutral mutations between two sequences that experience a reduction in recombination (Berdan et al. 2021). Divergence was stronger between arrangements of Cf-Inv(1) than between arrangements of the LG4 inversions. Several nonexclusive hypotheses can explain this. First, it is possible that Cf-Inv(1) is older, leaving more time for mutations to accumulate. Second, Cf-Inv(1) is a complex structural variant, which involved at least three separate inversion events (Aziz 1975; Day et al. 1982), and such complexity is known to suppress double crossovers and gene conversion, which maintain some exchange in simpler inversions (Korunes and Noor 2019). Finally, the distribution of karyotypes across the populations will strongly affect mutation accumulation by dictating the frequency of the arrangements (and thus their N_e) as well as the extent of recombination suppression. Cf-Inv(1) is polymorphic in all populations studied with a higher than expected proportion of heterokaryotypes. Conversely, Cf-Inv(4.1) has high frequencies of opposing homokaryotypes at each end of the cline. It is probably a

combination of age, extent of gene flux (i.e., double crossing over and gene conversion between arrangements), and karyotype distribution that explains the variation in differentiation between *C. frigida*'s inversions.

Chromosomal Inversions Are Involved in Adaptation to Heterogeneous Environments

Across geographic and ecological gradients, inversions may contribute strongly to genetic differentiation and often appear as islands of differentiation (Hoffmann et al. 2004). For instance, in the mosquito Anopheles gambiae, genetic differentiation along a latitudinal cline is almost entirely concentrated in two inversions (Cheng et al. 2012). In the marine snail Littorina saxatilis, genetic variation between habitats is largely driven by several inverted regions (Morales et al. 2019). Coelopa frigida follows this trend: pairwise F_{ST} values between populations are higher in inverted regions compared with collinear regions, albeit at a different geographic scale for the different inversions. Along the North-South gradient, differentiation between populations was higher and IBD was stronger in Cf-Inv(4.1) and Cf-Inv(4.2/4.3) than in collinear regions. F_{ST} based on SNPs in an inverted region combined two levels of genetic variation because differentiation between populations was driven by frequency variation at each highly differentiated arrangement. Such frequencies showed strong latitudinal clines, resembling the clines observed for several inversions in *Drosophila* that are maintained by selection-migration balance (Kapun et al. 2016). In sharp contrast, the genetic differentiation in the inverted region Cf-Inv(1) did not depend on geographic distances among populations. This pattern was related to the heterogeneous frequency of the α and β arrangements, which vary at a fine spatial scale but do not vary clinally. Yet, both the clines of Cf-Inv(4.1)/Cf-Inv(4.2/4.3) and the heterogeneity of Cf-Inv(1) contrasted with the homogeneous frequency of collinear variants, supporting the hypothesis that inversion distribution reflects spatial variation in selection pressures.

GEAs confirmed the putative role of inversions in adaptation to small-scale and large-scale variations of ecological conditions in *C. frigida*. Here, one question that may arise is whether the SNPs located in an inverted region are more likely to be detected as outliers than collinear SNPs. We avoided such artifacts by following the guidelines and best

practices from Lotterhos (2019) that used simulations to confirm the absence of bias when inversions or low-recombining regions were neutral. However, genome scan analyses are still more likely to detect adaptive regions with strong divergence that are resistant to swamping by migration, whereas dispersed, transient, or small-effect adaptive alleles are harder to detect (Yeaman 2015). Moreover, because of the high LD associated with an inversion, several SNPs may not be causative but simply linked to an adaptive variant. Hence, the high density of outlier SNPs in inverted regions neither means that they are full of adaptive alleles, nor that they are the only variants relevant for local adaptation. Nevertheless, the strengths of association between environment and the frequencies of some SNPs found in the inverted regions, as well as the association between environment and inversion frequencies, support inversions as major and true players of adaptation to heterogeneous environments in C. frigida. As such, the seaweed fly C. frigida joins an accumulating number of studies pioneered by Dobzhansky (1947, 1948), which have provided examples of species carrying several ecologically relevant inversions that are involved in local adaptation despite high gene flow (Anderson et al. 1991; Schaeffer 2008; Joron et al. 2011; Cheng et al. 2012; Kapun et al. 2016; Kirubakaran et al. 2016; Lindtke et al. 2017; Wellenreuther and Bernatchez 2018; Kapun and Flatt 2019; Huang et al. 2020). All of this is consistent with a model in which inversions are particularly relevant for adaptation with gene flow, because they preserve linkage between locally adapted alleles (Kirkpatrick and Barton 2006; Charlesworth and Barton 2018) and/or coadaptive epistatic alleles (Dobzhansky and Dobzhansky 1970; Charlesworth and Charlesworth 1973). However, although each inversion contains hundreds of genes, identifying multiple coselected or coadapted loci remains challenging because of LD, and calls for future experimental or transcriptomic work dissecting genetic variation in inversions.

In many empirical cases, when several inversions are found in the same species, they tend to vary along the same environmental axis. For instance, in the silverside fish Menidia menidia, several inverted haploblocks covary along a latitudinal gradient (Tigano et al. 2020; Wilder et al. 2020). The same tendency is observed for three inversions differentiating mountain and plain African honeybees Apis mellifera scutellata (Christmas et al. 2019), and dune and nondune ecotypes of the sunflower Helianthus petiolaris (Huang et al. 2020; Todesco et al. 2020). In contrast, for C. frigida, we observed two contrasting evolutionary patterns: The inversion Cf-Inv(1) was associated with wrackbed characteristics and composition, which represent patchy habitats at a fine geographic scale. It also functions as a supergene for body size, a trait which is usually polygenic yet appears in C. frigida to be controlled largely, if not entirely, by this inversion. The ecological and phenotypic associations are consistent with previous work on European and American populations (Day et al. 1983; Butlin and Day 1985; Berdan et al. 2018; Mérot et al. 2018). They reflect how the quality, composition, and depth of the wrackbed, possibly reflecting its stability, differently favor the opposite life-history strategies associated with the inversion. The β arrangement provides quick growth and

smaller size whereas the α arrangement provides high reproductive success linked to a larger size but at the expense of longer development time. This ecologically related tradeoff combined with heterozygote advantage results in strong balancing selection (Butlin 1983; Mérot, Llaurens, et al. 2020). Conversely, the inversions Cf-Inv(4.1) and Cf-Inv(4.2/4.3) show no deviation from Hardy-Weinberg disequilibrium and display a strong geographic structure along a latitudinal cline. As Cf-Inv(4.1) and Cf-Inv(4.2/4.3) are associated with climatic variables, we suggest that they possibly play a role in thermal adaptation. Additional support for this hypothesis come from the close proximity of Cf-Inv(4.1) and Cf-Inv(4.2/4.3) with a QTL for recovery after chill coma although we cannot exclude that the presence and the position of that QTL may suffer from mapping bias caused by low recombination (Noor et al. 2001). To summarize, these inversions describe contrasting patterns driven by different shapes of selection, with Cf-Inv(1) being a cosmopolitan polymorphism under balancing selection, whereas Cf-Inv(4.1) and Cf-Inv(4.2/4.3) represent geographically structured polymorphisms, possibly under spatially variable selection.

Exploring Low-Recombination Regions: What Are They and Why Do They Matter?

In addition to the aforementioned inversions, we also identified additional regions that spanned large fractions of each chromosome (6-16 MB) and were characterized by distinct haploblocks, high LD, and low recombination. With the current data, we can only speculate about what those regions are and what are the mechanisms underlying the observed patterns. Different types of data suggest different answers to this guestion. For example, the enrichment in transposable elements (supplementary fig. S7, Supplementary Material online) may indicate pericentromeric regions or transposonrich centromeres, which are challenging to assemble and characterize (Talbert and Henikoff 2020). However, we did not observe the typical enrichment of adénine and thymine(A/T)content (supplementary S7, Supplementary Material online). The landscape of nucleotide diversity was also very heterogeneous: parts of those lowrecombining regions are deserts of diversity (fig. 3C), as expected under increased selection at linked sites (also called "linked selection" [Cutter and Payseur 2013]), which leads to genetic hitchhiking around loci affected by positive or negative selection (Begun and Aquadro 1992; Charlesworth 1996). Yet, peaks of high diversity are observed in Cf-Lrr(2) and Cf-Lrr(5). These may reflect signatures of associative overdominance (Ohta 1971), due to masking of recessive deleterious loci in heterozygotes, as observed in some low-recombining regions of human and Drosophila genomes (Becher et al. 2020; Gilbert et al. 2020). Recent admixture from related lineages can also form distinct haploblocks (Li and Ralph 2019) and would generate similar patterns of high diversity but we consider this hypothesis unlikely in our case as no sympatric sister-species is known. Another possibility is that haploblocks coinciding with peaks of diversity are misassembled structural variants embedded in recombining region, such that haploblocks that are seemingly

separated could be adjacent. Our reference genome was scaffolded and ordered based on a linkage map from one family. Hence, inversions that were heterozygous in the mother, as well as any low-recombining regions, could cluster into large regions with low rates of crossing-over in the map, where marker ordering may be less accurate. Additional data such as long-reads or connected molecules like Hi-C are needed to improve the quality of the assembly in those specific areas and better characterize their DNA content. Despite these cautionary notes, our analysis provides an early annotation of regions that do not behave like the rest of the genome in terms of geographic genetic structure and association with environmental factors.

The low-recombining regions may also play a role in shaping the distribution of adaptive and nonadaptive variation since they differentiated populations more strongly than collinear regions. One potential reason would be increased variance in F_{ST} statistics in low-recombining regions that can emerge even under a purely neutral model (Booker et al. 2020). The effect of selection at linked sites, which reduces diversity in low-recombining regions, is also known to inflate differentiation, sometimes repeatedly between different pairs of populations (Burri et al. 2015; Hoban et al. 2016). Although these processes may explain extreme F_{ST} values in low diversity and low-recombining subregions, they are unlikely to explain the pattern that we observed in high-diversity subregions, at least in Cf-Lrr(2) and Cf-Lrr(5), which include GEA outliers as well as haplotypic variants whose frequency correlates with environmental variation. Without certainty about the mechanisms behind the reduced recombination, we can only propose hypotheses about the evolutionary processes at play. If these regions are complex or misassembled structural variants, they would represent additional adaptive chromosomal rearrangements contributing to adaptation in C. frigida, with different arrangements bearing one or several locally adapted alleles. If those regions are centromeric, or simply rarely recombining, they would highlight the importance of selection at linked sites in structuring intraspecific variation and the relevance of low-recombining regions in protecting locally adapted alleles. Evidence for an important evolutionary role of low-recombining regions is increasingly reported and we should analyze genomic landscapes in the light of recombination. For instance, in three-spine stickleback (Gasterosteus aculeatus), putatively adaptive alleles tend to occur more often in regions of low recombination in populations facing divergent selection pressures and high gene flow (Samuk et al. 2017). Similarly, regions of low recombination are enriched in loci involved in parallel adaptation to alpine habitat in the Brassicaceae Arabidopsis lyrata (Hämälä and Savolainen 2019). To what extent LD in low recombination regions affect such inferences yet remains an open question (Stevison and McGaugh 2020). Some statistics are biased by recombination heterogeneity (e.g., outliers based on F_{ST} in sliding-windows [Booker et al. 2020] or on PCA [Lotterhos 2019], QTL from mapping families [Noor et al. 2001]) but other approaches appear robust when following best practices (e.g., GEAs, selective sweep detection [Lotterhos 2019]). Overall, further work is needed, both on the methodological and empirical points of view, in order to better understand the contribution of recombination heterogeneity in structuring intraspecific variation and modulating migration-selection balance.

Conclusion

Our findings support the growing evidence that large chromosomal inversions play a major evolutionary role in some organisms characterized by extensive connectivity across a large geographical range. In this flying insect, as in several marine species, migratory birds, and widespread plants, chromosomal rearrangements strongly affect genetic diversity and represent a key component of the genetic architecture for adaptation in the face of gene flow. Critically, the different inversions are under different selective constraints across a range of geographical scales and contribute to adaptation to different environmental factors. Thus, inversions appear to be an architecture that allows some species to cope with gene flow as well as various sources and scales of environmental heterogeneity. Although inversions present one solution to the problem of adaptation with gene flow, it is still unknown how prevalent inversions are in nature. This is because structural variants are just beginning to be characterized in nonmodel species. In one of the best-studied clades, Drosophila, the answer is contradictory: closely related species D. melanogaster and D. simulans exhibit respectively more than 500 versus only 14 polymorphic inversions while both species are ecologically successful and distributed worldwide (Aulard et al. 2004). This is possibly due to the dichotomous nature of reduced recombination. Although reduced recombination holds together complexes of adaptive alleles, it also hampers the generation of new (and potentially adaptive) allele combinations and reduces the efficiency of purifying selection in linked regions (Felsenstein 1974). Overall, our analysis highlights the importance of regions of low recombination in structuring adaptive and nonadaptive intraspecific genetic variation. With recombination varying both along the genome and between individuals or haplotypes, inversions may represent only the simplest aspect of the complex relationship between recombination, selection, and gene flow that we are just starting to uncover through the prism of structural variants (Stapley et al. 2017). By optimizing wholegenome sequencing to include many individuals across a species range as done here, future work will have the possibility to better understand how the interplay between structural variation and recombination may matter for the evolution of biodiversity.

Materials and Methods

A Reference Genome Assembly for C. frigida

To generate a reference genome, we sequenced female siblings of C. frigida homozygous $\alpha\alpha$ for the inversion Cf-Inv(1), obtained by three generations of sib-mating from parents collected in St Irénée (QC, Canada). A pool of DNA from three siblings was sequenced on four cells of Pacific Biosystems Sequel sequencer, producing 16.1 Gbp (\sim 64x coverage) of long reads, and one sibling was sequenced with 10x Genomics Chromium on 1 lane of an Illumina HiSeqXTen

sequencer, yielding 82 Gbp (\sim 300x of coverage) of 150 bp paired-end linked-reads. Long reads were assembled using the Smrt Analysis v3.0 pbsmrtpipe workflow and FALCON (Chin et al. 2013), resulting in 2,959 contigs (N50 = 320 kb), for a total assembly size of 233.7 Mbp. This assembly was polished by using the linked-reads, first by correcting for sequence errors with Pilon (Walker et al. 2014) and, second, by correcting for misassemblies with Tigmint (Jackman et al. 2018). The resulting assembly consisted of 3,096 contigs (N50 = 320 kb). The contigs were scaffolded using the longrange information from linked-reads with ARKS-LINKS (Coombe et al. 2018), resulting in 2,539 scaffolds (N50 = 735 kb). Scaffolds were anchored and oriented into chromosomes using Chromonomer (Catchen et al. 2020), based on the order of markers in a linkage map (see below). The final assembly consisted of 6 chromosomes and 1,832 unanchored scaffolds (N50 = 37.7 Mb) for a total of 239.7 Mb (195.4 Mb into chromosomes). The completeness of this reference was assessed with BUSCO version 3.0.1 (Simão et al. 2015). The genome was annotated by mapping a transcriptome assembled from RNA sequences obtained from 8 adults (split by sex and by karyotype at the Cf-Inv(1) inversions), 4 pools of 3 larvae, and pools of C. frigida at different stages. The transcriptome was annotated using the Triannotate pipeline. More details are provided as supplementary methods, Supplementary Material online.

A High-Density Linkage Map and QTL Analyses Sequencing, Genotyping, and Building the Map

We generated an outbred F2 family of 136 progenies by crossing two F1 individuals of *C. frigida* obtained by crossing wild individuals collected in Gaspésie (QC, Canada). The mother of the F2 family was homozygous $\alpha\alpha$ at Cf-Inv(1). The progeny, both parents, and two paternal grandparents were sequenced using a double-digest restriction library preparation (ddRAD-seq) using ApeK1 on an IonProton (ThermoFisher), with greater depth for the parents. Reads were trimmed and aligned on the scaffolded assembly with bwa-mem. Genotype likelihoods were obtained with SAMtools mpileup. Only markers with at least 3X coverage in all individuals were kept. More details are provided in supplementary methods, Supplementary Material online.

A linkage map was built using Lep-MAP3 (Rastas 2017) following a pipeline available at https://github.com/clairemerot/lepmap3 pipeline (last accessed April 2021). Markers with more than 30% of missing data, noninformative markers, and markers with extreme segregation distortion (χ^2 test, P < 0.001) were excluded. Markers were assigned to LGs using the SeparateChromosomes module with a logarithm of odds (LOD) of 15, a minimum size of 5 and assuming no recombination in males, as is generally the case in Diptera (Satomura et al. 2019). This assigned 28,615 markers into 5 large LGs and 25 sex-linked markers into 2 small LGs than were subsequently merged as one (LG6). Within each LG, markers were ordered with five iterations of the OrderMarker module. The marker order from the run with the best likelihood was retained and refined three times with the evaluateOrder flag with five iterations each. When more than 1,000 markers were plateauing at the same position, all of them were removed, as suggested by *Lep-MAP* guidelines. Exploration for more stringent filtering or different values of LOD resulted in collinear maps.

Estimating Recombination Rate

To estimate recombination rate across the genome, we compared the positions of the markers along the genetic map with their position along the genome assembly with MAREYMAP (Rezvoy et al. 2007). Local recombination rates were estimated with a Loess method including 10% of the markers for fitting the local polynomial curve.

QTL Analysis

All individuals used to build the map were scored for recovery at room temperature after a chill coma induced by holding them for 10 min at $-20\,^{\circ}$ C. We distinguished three categories: "0," the fly stands immediately or in less than 5 min; "1," the fly recovers with difficulty after 5–15 min; "2," the fly has not recovered after more than 15 min. A phased map was obtained by performing an additional iteration of the *OrderMarker* module. QTL analysis was carried out using R/qtl (Broman et al. 2003). LOD scores correspond to the $-\log_{10}$ of the associated probabilities between genotype and phenotype with the Haley–Knott method. The LOD threshold for significance was calculated using 1,000 permutations.

Population-Level Sequencing

Sampling and Characterization of Size and Karyotype

We analyzed 1,446 adult *C. frigida*, sampled at 16 locations spanning over 10° of latitude (fig. 1A) in September/October 2016. Sampling, genotyping, and phenotyping are described in detail in Mérot et al. (2018). Size was estimated using wing length as a proxy for 1,426 flies. Genomic DNA was extracted using a salt extraction protocol (Aljanabi and Martinez 1997) with an RNase A treatment (Qiagen). A total of 1,438 flies were successfully genotyped at the *Cf-Inv*(1) inversion using a molecular marker (Mérot et al. 2018).

Library Preparation, Sequencing, and Processing of the Sequences

Whole-genome high-quality libraries were prepared for each fly by adapting a protocol described in Baym et al. (2015) and Therkildsen and Palumbi (2017) and detailed in supplementary materials, Supplementary Material online. Briefly, DNA extracts were quantified, distributed in 17 plates with randomization (96-well), and normalized at 1 ng/µl. Each sample extract underwent a step of tagmentation, which fragments DNA and incorporates partial adapters, two PCR steps that attached barcode sequences (384 combinations) while amplifying the libraries, and a size selection step using an Axygen magnetic bead cleaning protocol. Final concentrations and fragment size distributions were estimated to combine equimolar amounts of 293 to 296 libraries into five separate pools. Sequencing on five lanes of Illumina HiSeq 4000 yielded an

average of 327 Mb per sample, which resulted in approximately 1.4X coverage (range: 121–835 Mb, 0.5X-3.5X).

Paired-end 150-bp reads were trimmed and filtered for quality with FastP (Chen et al. 2018), aligned to the reference genome with BWA-MEM (Li and Durbin 2009), and filtered to keep mapping quality over 10 with Samtools v1.8 (Li et al. 2009). Duplicate reads were removed with MarkDuplicates (PicardTools v1.119.) We realigned around indels with GATK IndelRealigner (McKenna et al. 2010) and soft clipped overlapping read ends using clipOverlap in bamUtil v1.0.14 (Breese and Liu 2013). Reads, in bam format, were analyzed with the program ANGSD v0.931 (Korneliussen et al. 2014), which accounts for genotype uncertainty and is appropriate for low-coverage whole-genome sequencing (Lou et al. 2020). Input reads were filtered to remove low-quality reads and to keep mapping quality above 30 and base quality above 20.

As a first step, we ran ANGSD to estimate the spectrum of allele frequency, minor allele frequency (MAF), depth, and genotype likelihoods on the whole data set. Genotype likelihoods were estimated with the GATK method (-GL 2). The major allele was the most frequent allele (-doMajorMinor 1). We filtered to keep positions covered by at least one read in at least 50% of individuals, with a total coverage below 4,338 (three times the number of individuals) to avoid including repeated regions in the analysis. From this list of variant and invariant positions, we selected SNPs with an MAF of above 5% and subsequently used this list with their respective major and minor alleles for most analyses (PCA, inversion detection, F_{ST}, GEAs). Using PLINK 1.9, we selected a subset of SNPs pruned for physical linkage, removing SNPs with a variance inflation factor greater than two (VIF > 2) in 100 SNP sliding windows shifted by five SNPs after each iteration.

PCA and Inversion Detection

An individual covariance matrix was extracted from the genotype likelihoods in beagle format using PCAngsd (Meisner and Albrechtsen 2018) and decomposed into PCs with R, using a scaling 2 transformation, which adds an eigenvalue correction (Legendre and Legendre 2012). To analyze genetic variation along the genome, we performed the same decomposition in nonoverlapping windows of 100 SNPs. For each "local PCA," we analyzed the correlation between PC1 scores and PC scores from the PCA performed on the whole genome. This allowed us to locate two (inversion) regions underlying the structure observed on PC1 and PC2 (supplementary fig. 2A, Supplementary Material online). We set the boundaries of those regions as windows with a coefficient of correlation above one standard deviation.

To scan the genome for other putative inversions or non-recombining haploblocks, we used the R package *Lostruct* (Li and Ralph 2019), which measures the similarity between local PCA (PC1 and PC2 for each 100 SNP window) using Euclidean distances. Similarity was mapped using MDS of up to 20 axes. Clusters of outlier windows (presenting similar PCA patterns) were defined along each MDS axis as those with values

beyond 4 standard deviations from the mean, following Huang et al. (2020). Adjacent clusters with less than 20 windows between them were pooled, and clusters with less than 5 windows were not considered. Different window sizes (100 to 1,000), different subset of PCs (1 to 3 PCs), and different thresholds yielded consistent results. A typical signature of a polymorphic inversion is three groups of individuals appearing on a PCA: the two homokaryotypes for the alternative arrangements and, as an intermediate group, the heterokaryotypes. All clusters of outlier windows were thus examined either by a PCA as single blocks, or divided into several blocks when discontinuous. We then used K-means clustering to identify putative groups of haplotypes. Clustering accuracy was maximized by rotation and the discreteness was evaluated by the proportion of the between-cluster sum of squares over the total.

Inversion Analysis

For the four inversions (Cf-Inv(1), Cf-Inv(4.1), Cf-Inv(4.2/4.3)), K-means assignment on PC scores was used as the karyotype of the sample. Differentiation among karyotypes was measured with F_{ST} statistics, using ANGSD to estimate joint allele frequency spectrum, realSFS functions to compute F_{ST} in sliding windows of 25 KB with a step of 5 KB, and subsampling the largest groups to balance sample size. Observed proportion of heterozygotes (Hobs) was calculated for each karyotype and each SNP using the function -doHWE in ANGSD, and then averaged across sliding windows of 25 KB with a step of 5 KB using the R package windowscanr. Nucleotide diversity (π) within each arrangement, and nucleotide divergence (dxy) between arrangements was calculated in sliding windows of 25 KB (step 5 KB) considering all positions (variants and invariants), controlling for missing positions and using the function doThetas (ANGSD) and the script https://github.com/mfumagalli/ngsPopGen/blob/master/scripts/calcDxv.R accessed April 2021), following the recommendation of Korunes and Samuk (2021). For each 25-kb window, nucleotide divergence was corrected for within arrangement genetic variation by subtracting the mean of the nucleotide diversity in both arrangements. The 95% confidence intervals were estimated by bootstrapping the data using perwindow corrected d_{XY} estimates (1,000 replicates). Based on this value of corrected d_{XY} between each inversion's arrangements calculated on noncoding windows in inverted regions, we estimated an approximate time of divergence using a constant molecular clock. We assumed a mutation rate comparable to Drosophila, with μ equal to 5×10^{-9} mutations per base per generation (Assaf et al. 2017). Given that generation time in C. frigida strongly varies, from 8 to 20 days at 25 °C up to months in colder conditions, we considered a range of five to ten generations per year. As arrangements are expected to keep some gene flux after the formation of the inversion, due to double crossing-over and gene conversion (Navarro et al. 1997; Korunes and Noor 2019), the age estimates should be considered as a minimum value.

Linkage Disequilibrium

Intrachromosomal LD was calculated among a reduced number of SNPs, filtered with more stringent criteria (MAF > 10%, at least one read in 75% of the samples). Pairwise R^2 values were calculated with NGS-LD (Fox et al. 2019) based on genotype likelihood obtained by ANGSD, and grouped into windows of 1 MB. Plots display the 2nd percentile of R^2 values per pair of windows. For LG1 and LG4, R^2 was calculated, first within all samples, then within individuals homozygous for the most common orientation of each inversion, subsampling the largest groups to balance sample size, and plotted by windows of 250 kb.

Geographic Structure

 F_{ST} differentiation between all pairs of populations was estimated based on the allele frequency spectrum per population (-doSaf) and using the realSFS function in ANGSD. Positions were restricted to the polymorphic SNPs (>5% MAF) previously polarized as major or minor allele (options –sites and – doMajorMinor 3), and which were covered in at least 50% of the samples in a given population. Populations were randomly subsampled to a similar size of 88 individuals. The weighted F_{ST} values between pairs of population were computed by including either all SNPs, LD-pruned SNPs, or SNPs from a region of interest (inversions/low-recombining regions) or SNPs outside those regions (collinear SNPs).

IBD was tested for each subset of SNPs using a linear model in which pairwise genetic distance $(F_{ST}/(1-F_{ST}))$ was included as the response variable and geographic Euclidean distance was incorporated as an explanatory term. IBR refers to constrained dispersal due to environmental features that limit movement and was tested in the same way as IBD, except that physical distances were calculated along the shoreline, assuming that the open water or mainland may oppose dispersal of C. frigida. The distance via least cost path was measured through areas of the map between -40 m of depth and 20 m of altitude using the R package marmap. Both models of IBD and IBR were compared with a null model using an ANOVA F-test, and to each other using adjusted R² and AIC (Akaike Information Criteria). To compare IBD and IBR patterns in each inversion/low-recombining region to the collinear genome, we built a full model explaining pairwise genetic distances by physical distances and genomic region (collinear vs. inversion) as a cofactor, and assessed the significance of the interaction term as well as the direction of the interaction slope coefficient. We repeated this analysis 100 times with randomly chosen collinear regions including the same number of contiguous SNPs as each inversion/lowrecombining regions. This provided a distribution of the significance of the interaction term and its slope coefficient (supplementary fig. S13, Supplementary Material online). For Cf-Inv(1), no contiguous block with the same number of SNPs could be found in the genome, hence we gathered 3 blocks of 1/3 the number of SNPs in each of the 100 random replicates.

Finally, we examined the direct association between inversions and latitude, treating inversions as single bi-allelic loci.

The association was tested by a Generalized Linear Model (GLM) with a logistic link function for binomial data, the response variable being the number of individuals carrying/ not carrying the minor arrangement and the explanatory variable being latitude. To assess whether this association deviates from null expectations, we randomly sampled 1,000 SNPs, with an average frequency similar to each inversion, and built 1,000 full models explaining frequency by latitude and genomic region (a collinear SNP vs. an inversion) as a cofactor, and assessed the significance of the interaction term (supplementary fig. S15, Supplementary Material online).

Environmental Associations

Environment at each location was described by large-scale climatic/abiotic conditions and local wrackbed characteristics (supplementary table S1, Supplementary Material online), as described in Mérot et al. (2018). Large-scale conditions included annual means in precipitation, air temperature, sea surface temperature, sea surface salinity, and tidal amplitude extracted from public databases. Wrackbed characteristics were measured upon collection and included abiotic variable (depth, internal temperature, and salinity) and algal composition (relative proportions of Laminariaceae, Fucaceae, Zoosteraceae, plant debris, and other seaweed species). Correlation between variables was tested with a Pearson correlation test, and variation was reduced by performing a PCA for each group of correlated environmental variables (climatic, salinity/tidal amplitude, abiotic characteristic of the wrackbed, algal composition) and retaining significant PCs following the Kaiser-Guttman and Broken Stick criteria (Borcard et al. 2011) (see supplementary fig. Supplementary Material online).

MAF was calculated for each population from the list of SNPs previously polarized as major or minor allele (–sites and – doMajorMinor 3), and covered by at least 50% of the individuals in each population. Allelic frequency was thus estimated with confidence (>50X of coverage at population level) for a total of 1,155,978 SNPs. A GEA that evaluated SNPs frequencies as function of environmental variables was performed through a combination of two methods as recommended by de Villemereuil et al. (2014): 1) latent factor mixed models (LFMM2; Frichot et al. 2013; Caye et al. 2019) and 2) Bayes factor (BAYPASS; Gautier 2015). Those two methods had also been shown to be robust to the presence of large inversions (Lotterhos 2019).

LFMM was run with the R package *lfmm2* (Caye et al. 2019) on the full set of 1,155,978 SNPs, using a ridge regression, which performed better in simulations including inversions (Lotterhos 2019), and parametrized using a K-value of four latent factors (as evaluated from a PCA on an LD-pruned data set). False discovery rate (FDR) was assessed following the recommendations of François et al. (2016), using a Benjamini–Hochberg correction. Using Baypass v2.2 (Gautier 2015), a BF, evaluating the strength of an association between SNP frequency and environment, was computed as the median of three runs under the standard model on the full set of 1,155,978 SNPs. Environmental variables were scaled using the

-scalecov option. We ran this analysis twice: first, without controlling for population structure and, second, by controlling with a covariance matrix extracted from an initial BayPass model run on the subset of LD-pruned SNPs without environmental covariables. To calculate a significance threshold for BF, we simulated pseudo-observed data with 10,000 SNPs using the "simulate.baypass" function and kept the 0.1% quantile as the significance threshold. For each GEA method, and the combination of the two, the repartition of candidate SNPs for association with environment inside and outside inversions/low-recombining regions was compared with the original repartition of SNPs. Deviation from this original repartition was tested with a Fisher's exact test, and the magnitude of the excess/deficit of outlier SNPs in each region of the genome was reported as the odd ratio.

We also compared the distribution of association scores in each inversion/low-recombining region to the collinear genome. This test was performed on absolute values of the z scores from LFMM, using a generalized linear model with quasinormal family (square root link) and genomic region (collinear vs. inversion) as an explanatory factor. We repeated this analysis on 100 randomly chosen collinear blocks including the same number of SNPs as each inversion/low-recombining regions times (supplementary fig. S19, Supplementary Material online). Finally, we examined the direct association between inversions and environment variables, treating each inversion as a single locus, as described above for latitude using GLM models and comparing to 1,000 randomly drawn SNPs (supplementary fig. S20, Supplementary Material online).

Phenotypic Associations and Gene Ontology Analysis

We performed a GWAS for wing size using ANGSD latent genotype model (EM algorithm, -doAssso = 4) where genotype is introduced as a latent variable and then the likelihood is maximized using weighted least squares regression (Jørsboe and Albrechtsen 2020). We considered a FDR of 0.001. The GWAS was applied on the whole data set (1,426 flies with size information) and then on each subset of homokaryotypes at the inversion *Cf-Inv*(1) (140 $\alpha\alpha$ and 436 $\beta\beta$ flies with size information).

Using BEDtools, we extracted the list of genes overlapping with significantly associated SNPs, or within a window of 5 kb upstream or downstream a gene. We then tested for the presence of overrepresented GO terms using GOAtools (v0.6.1, $P_{\rm val}=0.05$) and filtered the outputs of GOAtools to keep only GO terms for biological processes of levels 3 or more, and with an FDR value equal below 0.1. We performed the same GO enrichment analysis for the list of genes found in the two largest inversions (*Cf-Inv*(1) and *Cf-Inv*(4.1)).

Supplementary Material

Supplementary data are available at Molecular Biology and Evolution online.

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Data Availability

The genome assembly, GBS reads used to build the linkage map and WGS paired reads used for population genomics are available on NCBI under the projects, respectively (PRJNA688905, PRJNA689789, and PRJNA689963). Raw information about GBS and WGS samples is provided as supplementary tables S9 and S10, Supplementary Material online. The pipelines for analyses of WGS data are available at https://github.com/enormandeau/wgs_sample_preparation (last accessed April 2021) and https://github.com/clairemerot/angsd_pipeline (last accessed April 2021).

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