Cold adaptation drives population genomic divergence in the ecological specialist, *Drosophila montana*

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Abstract
Detecting signatures of ecological adaptation in comparative genomics is challenging, but analysing population samples with characterised geographic distributions, such as clinal variation, can help identify genes showing covariation with important ecological variation. Here, we analysed patterns of geographic variation in the cold-adapted species *Drosophila montana* across phenotypes, genotypes and environmental conditions and tested for signatures of cold adaptation in population genomic divergence. We first derived the climatic variables associated with the geographic distribution of 24 populations across two continents to trace the scale of environmental variation experienced by the species, and measured variation in the cold tolerance of the flies of six populations from different geographic contexts. We then performed pooled whole genome sequencing of these six populations, and used Bayesian methods to identify SNPs where genetic differentiation is associated with both climatic variables and the population phenotypic measurements, while controlling for effects of demography and population structure. The top candidate SNPs were enriched on the X and fourth chromosomes, and they also lay near genes implicated in other studies of cold tolerance and population divergence in this species and its close relatives. We conclude that ecological adaptation has contributed to the divergence of *D. montana* populations throughout the genome and in particular on the X and fourth chromosomes, which also showed highest interpopulation *F*ₜ. This study demonstrates that ecological selection can drive genomic divergence at different scales, from candidate genes to chromosome-wide effects.

Keywords
chill coma recovery time, cline populations, cold tolerance, CTmin, *D. montana*, environmental adaptation, genomic divergence
INTRODUCTION

The geographic structure of a species is a result of its phylogeographic history, influenced by past and present dispersal, population demography, and selection. Obtaining genome-wide data on genetic polymorphisms across multiple populations of a species is becoming relatively easy, but interpreting the patterns of geographic variation in such data and identifying genes which vary primarily due to selection remains challenging. Often simple “outlier” approaches using genome scans which measure genetic differentiation such as $F_{ST}$ or Dxy are adopted, but results are difficult to interpret due to confounding effects of selection, drift and population structure, or genomic features such as inversions and other causes of variation in recombination rate (Cruickshank & Hahn, 2014; Noor & Bennett, 2009; Ravinet et al., 2017; Wolf & Ellegren, 2016). If environmental data are available, we can use associations with such factors to help identify loci where genetic differentiation covaries with this environmental variation. Some genome scan methods can incorporate environmental variation and simultaneously fit effects for covariance with environmental factors, while controlling for effects of population demography (Foll & Gaggiotti, 2008; de Villemereuil & Gaggiotti, 2015). This approach has successfully identified genetic variation associated with altitude in humans, among other examples (Foll et al., 2014; Gautier, 2015; de Villemereuil & Gaggiotti, 2015) and has become a useful approach to investigate the ecological adaptations underlying population divergence.

Clinal patterns of variation in phenotypes or gene frequencies have a long history of being used to infer selection along ecotones, and analyses ofcline shape can sometimes identify loci under direct selection from others showing clinal variation for other reasons, such as phylogeographic history (Barton & Gale, 1993). Such studies can be very powerful, especially when independent parallel clines are available. For example, Kolaczkowski et al., (2011) sampled isofemale lines from extremes of a cline in Australian populations of D. melanogaster, and found many genes implicated in clinally varying phenotypes to show highest differentiation. Also, Bergland et al., (2014), and Kapun et al., (2016) sampled North American clines in D. melanogaster and D. simulans over several years to uncover clinal variation at a genome scale. Bergland et al., (2014) also found consistent fluctuations in allele frequencies for a population sampled over several seasons, indicating a regular response to seasonally varying selection pressures. On the other hand, Machado et al., (2015) concluded that migration and gene flow play a greater role than adaptation in the overall clinality of genomic variants in D. simulans than D. melanogaster. While the two species share a significant proportion of the genes showing clinal variation, their differences in overwintering ability, migration and population bottlenecks probably act as additional drivers of differences in patterns of variation between them (Machado et al., 2015). Similar studies of clinal variation in phenotypes and allele frequencies have also been carried out in other insects (Paolucci et al., 2016), plants (Bradbury et al., 2013; Chen et al., 2012), mammals (Carneiro et al., 2013; Hoekstra et al., 2004), fish (Vines et al., 2016), and other organisms (Endler, 1973, 1977; Takahashi, 2015). However, more studies are needed that simultaneously compare patterns of variation in allele frequencies and potentially causal environmental variation or ecologically important traits while controlling for population structure, as such studies are necessary to determine if adaptation to climate is directly driving patterns of genetic differentiation.

Here, we investigated geographic variation at both the phenotypic and genetic levels in Drosophila montana samples from two continents. This species has spread around the northern hemisphere (Throckmorton, 1969) and both mtDNA and microsatellite data have revealed genetically distinct Finnish and North American populations (Mirol et al., 2007). Moreover, more recent modelling of genome-wide SNP frequencies suggest that the Finnish-North American split happened around 1.75 Mya and that of the North American populations shortly after that (Garlovsky et al., 2020). It is one of the most cold-tolerant Drosophila species (Kellermann et al., 2012) and the basic cold tolerance of D. montana flies can increase towards the cold seasons through two mechanisms, photoperiodic reproductive diapause (Vesala & Hoikkala, 2011) and cold-acclimation induced by a decrease in day length and/or temperature (Kauranen et al., 2019; Vesala, Salminen, Kostal, et al., 2012; Vesala, Salminen, Laiho, et al., 2012). D. montana populations have been found to show clinal variation in the critical day length required for diapause induction (CDL; Lankinen et al., 2013; Tyukmaeva et al., 2011). There is also a correlation between CDL and latitudinally covarying climatic factors such as the mean temperature of the coldest month (Tyukmaeva et al., 2020). In addition, D. montana populations from different geographic regions show variation in their courtship cues and mate choice (Klappert et al., 2007; Routtu et al., 2007), which has led to partial reproductive isolation between some distant populations (Jennings et al., 2011, 2014). At the genetic level, differential gene expression studies have identified candidate genes underlying diapause (Kankare et al., 2010, 2016), perception of day length (Parker et al., 2015). Furthermore, a quasi-natural selection experiment for shorter CDL, accompanied by a decrease in cold-tolerance, induced widespread changes in loci with potential roles with these traits (Kauranen et al., 2019). Finally, population genomic analyses have identified several outlier loci when examining differentiation between North American and European populations (Parker et al., 2018). All this makes D. montana an interesting example of nascent speciation, potentially influenced by ecological adaptation.

Here, we sought to ask to what extent patterns in the genomic divergence of D. montana populations across two continents are correlated with climatic variation and phenotypic responses to cold adaptation. We performed pooled whole-genome sequencing (pool-seq) on six different populations and used Bayesian methods to examine the association between genomic differentiation between populations and environmental variables across both continents. We also phenotyped populations for two different cold tolerance measures; critical thermal minimum (CTmin), and chill coma recovery time (CCRT), and investigated the associations between them and the genetic and climatic data. Ultimately, we
asked if the genomic loci showing an association between genetic and environmental differentiation also showed association with population differentiation in cold tolerance phenotypes, and examined the possible overlap between the set of genes close to candidate SNPs with sets of candidate genes from previous studies of cold adaptation in \textit{D. montana}. If population differentiation is driven by ecological selection then we would predict the extreme cold adaptation of \textit{D. montana} to have left a signature of genomic divergence associated with environmental and phenotypic differentiation across these loci.

2 | MATERIALS AND METHODS

2.1 | Sample collections and DNA extraction

We collected samples of 49–50 wild-caught flies from six \textit{D. montana} populations from a range of latitudes from 66°N to 38°N in the spring of 2013 or 2014. Four of these populations represented a range of latitudes in North America (N.A.), and two populations were from a range in Finland (Figure 1; Table 1). Samples of wild-caught flies from the six populations were stored in ethanol (the male/female ratio varied across samples; Table 1) and DNA of individual flies was extracted using CTAB solution and phenol-chloroform-isoamylalcohol purifications in 2016. Genomic DNA was extracted from individual flies and quantified using Qubit (Thermo Fisher Scientific), and an equal amount of DNA from each individual (50 ng) was pooled into the final sample. Sequencing was performed at the Finnish Functional Genomics Centre in Turku, Finland (www.btk.fi/functional-genomics) on the Illumina HiSeq3000 platform (paired-end reads, read length = 150 bp, estimated coverage ~121x).

2.2 | Phenotyping

We measured the critical thermal minimum (CTmin) and chill coma recovery time (CCRT) of flies from six populations. Fly samples for these tests were collected for five populations (Seward, Terrace, Ashford, Crested Butte, and Korpilahti) from mass-bred populations that have been maintained in the laboratory since 2013–2014. For the Oulanka population, flies were collected from three isofemale strains (established in 2014), because the mass-bred population had been contaminated by another species. The mass-bred populations were originally established from F4 progenies of 20 isofemale strains each (400 flies) and have been maintained in constant temperature (19 ± 1°C) and light (24 h of light, to prevent the flies from entering diapause) regimes for about 20–25 generations before the experiment. All flies were supplied with fresh malt medium in half-pint bottles every week (Lakovaara, 1969). The newly emerged flies were collected using light CO₂ anaesthesia within 24 h after emergence, separated by sex and placed in malt-vials in the same conditions until sexual maturity (20–21 days old) and were then used in CTmin and CCRT tests. The same individual flies were first assayed for CTmin and then for CCRT, the flies were not anaesthetised before these tests.

We assayed a total of 328 females and 302 males for CTmin and CCRT. These assays were done in batches of between 22 and 30 flies, and split evenly by sex (21 batches in total). Between 32 and 46 (mean 39) flies per population per sex were tested (for the
TABLE 1 The sources of genomic samples (coordinates and the name of the nearest town), altitude of the sampling site, the year in which sampling was performed, and the number of males and females sampled (M/F) for each pool

| Source              | Sampling site                  | Year | M/F |
|---------------------|-------------------------------|------|-----|
| USA, Alaska         | Seward 60°9′ N; 149°27′ W    | 2013 | 30/20 |
| Canada, British Columbia | Terrace 54°27′ N; 128°34′ W | 2014 | 22/27 |
| USA, Washington     | Ashford 46°45′ N; 121°57′ W   | 2013 | 16/34 |
| USA, Colorado       | Crested Butte 38°54′ N; 106°57′ W | 2013 | 36/13 |
| Finland             | Oulanka 66°22′ N; 29°20′ E    | 2013 | 25/25 |
| Finland             | Korpihahti 62°20′ N; 25°34′ E | 2013 | 27/23 |

Oulanka isofemale strains, between 15 and 39 flies per strain per sex were used). CTmin tests are based on detecting the temperature (CTmin) at which flies lose neuromuscular function and enter reversible state of chill coma (Andersen et al., 2015). In these tests, the flies were placed into tubes sealed with parafilm and submerged into a 30% glycol-water mixture in Julabo F32-HL chamber. The temperature was decreased at the rate of 0.5°C per min (from 19°C to −6°C) and CTmin was determined by eye, as the temperature at which a fly was unable to stand on its legs. Immediately after the CTmin test, the temperature was set to −6°C and the flies were left in this temperature for 16 h. Vials were then quickly taken out of the glycol-water bath and the flies’ CCRT was determined as the time required for the flies to recover from chill coma and stand on their legs. The ambient room temperatures was recorded during trials but in an initial analysis including source population and room temperature there was no statistically significant effect of room temperature on CCRT (F_{5,569} = 0.72, p = .4), this variable was therefore left out of further analyses.

To investigate population differences in CTmin and CCRT phenotypes we fit general additive mixed models (GAMMs) in R (v. 3.6.3; R Development Core Team, 2020) using the “mgcv” R package (Wood, 2004). In simple linear models including population, sex, and experimental batch as a fixed effects, experimental batch had an effect on CTmin (F_{20,533} = 2.8, p < .01), although compared to population (F_{5,533} = 12, p < .01), and sex (F_{1,533} = 11.5, p < .01) this effect was not strong. Experimental batch had a stronger effect on CCRT (F_{20,533} = 2.83, p < .01), compared to population (F_{5,533} = 4.5, p < .01) and sex (F_{1,533} = 1.5, p < .22). We therefore included it as a random effect in GAMM analyses. The full GAMMs included altitude, latitude, and sex as fixed effects and experimental batch as a random effect. We used a cubic regression spline as the basis smoothing function for both altitude and latitude. The raw data for all phenotyping are given in Table S1. The full models are shown in the Supporting Information (Supplementary Material).

2.3 | Bioclimatic variables and population geography

We obtained representative climate data from the WorldClim database (Hijmans et al., 2005) for each D. montana population sampled for the pool-seq (see above), as well as for 18 additional populations of this species (Table S1; Tyukmaeva et al., 2020). We downloaded the climate data and extracted the values corresponding to population coordinates using the R package “raster” (version 2.5-8; Hijmans et al., 2016). In total this amounts to 55 bioclimatic variables for each population (Table S1). To reduce the number of variables in the data set a principle components analysis (PCA) was performed using the “PCA()” function from the R package “FactoMineR” (version 1.28; Lé et al., 2008). Principle components were kept for further analysis if their eigenvalues were >1. PCA scores for each population were z-transformed using the “scale()” function in base R. Additionally, CTmin and CCRT were summarised to a mean value for each population. In total, this gives four “environmental” variables measured for each population (PC1, PC2, CTmin, and CCRT).

2.4 | Mapping, SNP calling and genomic analysis

Quality of raw reads was checked with fastqc (v. 0.11.5; Andrews, 2015) and reads were trimmed using trimmomatic (v. 0.32; Bolger et al., 2014; see Supporting Information (Supplementary Material) for full trimming parameters). Trimmed reads were mapped to the D. montana reference genome (Parker et al., 2018) using bwa mem (v. 0.7.7; Li, 2013) with the default options but keeping only alignments with a mapping quality of >20 following best practice guidelines for pool-seq (Schlötterer et al., 2014). Duplicate alignments were removed with samtools rmdup (v. 1.3.1; Li et al., 2009) and regions around indels were realigned using picard (version 1.118; Broad Institute), gatk (v. 3.2-2; McKenna et al., 2010) and samtools. Separate.bam files for each of the sequenced populations were finally merged using bamtools (v. 2.4.0; Barnett et al., 2011).

Over 80% of reads were properly mapped and retained in all samples. The mean coverage for Seward samples was nearly twice that of the other samples (Figure S1 and Figure S2). To remove the potential for this difference to cause artefacts in downstream analyses, the.bam files for Seward were downsampled to contain 94.1 million reads (the average across the five remaining populations). Median empirical coverage was between ~62 and 88x (Table S2; Figure S3) and much less variable among the populations, allowing common maximum and minimum thresholds to be set based on the aggregate distribution. Allele counts for each population at each genomic position were obtained with samtools mpileup (version 1.3.1; Li et al.,
2009) using options to skip indel calling as well as ignoring reads with a mapping quality <20 and sites with a base quality <15. This was followed by the heuristic SNP calling software PoolSNP using a minimum count of 5 to call an allele, and a minimum coverage of 37 and a maximum coverage <95th percentile of the scaffold-wide coverage distribution to call a SNP (Kapun et al., 2020). Even if all these filters were passed, an allele was not considered if its frequency was <0.001. Finally, we only considered SNPs on scaffolds >10 kb in length. The final set consisted of 2,190,511 biallelic SNPs that could be placed on scaffolds ordered along the chromosomes and were used in downstream analyses.

To test for an association between the four environmental variables and genetic differentiation we used BAYESCOENV (v. 1.1; Foll & Gaggiotti, 2008; de Villemereuil & Gaggiotti, 2015). BAYESCOENV fits a model of \( F_{ST} \) to population differentiation for each locus, incorporating environmental differentiation as a predictor (included as the parameter \( g \)) while fitting two locus-specific effects, one for environmental selection, and the other for other processes (demography, other types of selection). It therefore controls for confounding effects of population structure/relatedness in testing for an association with environmental variables. BAYESCOENV was run with five pilot runs of 1000 iterations each, followed by a main chain of 4000 iterations of which 2000 were discarded as burnin. Four MCMC chains were run for each analysis to evaluate convergence of the chains to common parameter estimates. Because of the unbalanced number of males (and the resulting variation in ratios of X:Y chromosomes) in the pools, BAYESCOENV analyses were performed separately on SNPs that could be assigned to the autosomal linkage groups (chromosomes) and the X chromosome. Raw count data were used for the autosomal data. For X linked SNPs, allele count data were scaled to the known number of X chromosomes in the pool using \( n_{eff} \), the effective sample size taking into account the multiple rounds of binomial sampling inherent to a pool-seq design (Feder et al., 2012; Kolaczkowski et al., 2011). \( n_{eff} \) scales the allele counts at each SNP downwards based on the known number of chromosomes in the pool (see Table 1).

Chains were assessed for convergence with the “coda” \* package (v. 0.19-1; Plummer et al., 2006). Convergence was reached across the four chains for most analyses and parameters (potential scale reduction factors (PSRFs) of ~1 in a Gelman-Rubin diagnostic test; Figures S4–S7), except for analyses of autosomal SNPs and PC2 as the environmental variable which showed mild convergence problems (PSRF = 1.71), although parameter estimates agreed well with the other chains. Thus, this first chain (black line in Figures S4–S7) was discounted for all parameters and only estimates from the remaining three chains were used. The union of significant SNPs (using \( q \)-values for the \( g \) parameter, describing the association between genetic differentiation at a locus and environmental differentiation, to control the FDR at 0.05, i.e., SNPs with \( q \)-values <0.05) across these chains were taken as the final candidate SNPs.

Finally, population genetic statistics (\( x \), and Tajima’s \( D \)) were computed in windows of 10 kb with a step size of 5 kb using methods implemented for pool-seq data (Kapun et al., 2020). Windows contained a mean of 488.2 ± 1.2 SNPs (mean ± SE). These statistics were only computed for scaffolds with a length >50 kb. SNP-wise \( F_{ST} \) was computed for each population with the \( \tau \) package “poolfstat” (v. 1.1.1; Hivert et al., 2018) by first computing all pairwise values, and then deriving population specific \( F_{ST} \) values by averaging across all pairwise values where a population was included. We computed 95% confidence intervals for mean SNP-wise \( F_{ST} \) on each chromosome from the distribution of mean \( F_{ST} \) values across 100 bootstrap samples of SNPs. See the Supplementary Material (Supporting Information) for pseudocode commands of the key pipeline steps.

We performed GO term enrichment analyses with DAVID (v. 6.8; Huang et al., 2009a, 2009b) and GOWINDA (v. 1.12; Kofler & Schlötterer, 2012), which accounts for differences in gene length. For DAVID, since the \( D. montana \) annotation contains information about orthologs in \( D. virilis \), we extracted all genes within 10 kb upstream or downstream of a candidate SNP with an ortholog in \( D. virilis \) and submitted them to DAVID. For analyses with GOWINDA, because the gene-sets need to be given manually, we obtained gene-sets from FuncAssociate3 (Berriz et al., 2009). As above, we considered SNPs within 10 kb of a gene (~gene-definition updownstream 10,000) and performed 1 million simulations to obtain the empirical \( p \)-value distribution. Regulatory elements, such as enhancers, and transcription factor binding sites, can occur up to 1 Mb up- or downstream from a target gene in other species (Chan et al., 2010; Maston et al., 2006; Pennacchio et al., 2013; Werner et al., 2010) but generally lie within 2 kb of a gene region in \( D. melanogaster \) (Arnosti, 2003), thus 10 kb represents a compromise.

## 3 | RESULTS

### 3.1 | Cold tolerance measures, bioclimatic variables and population geography

Across individuals, there was no evidence of an association between minimum critical temperature (\( CT_{min} \); pooled across sexes) and the chill coma recovery time (CCRT; \( p = 0.08, p = .07 \)). Neither was there evidence for an association between these traits across populations (\( p = 0.22, p = .68 \)). \( CT_{min} \) was significantly different between sexes, with females being more cold tolerant (\( t = 3.14, p = .002 \)). While latitude appears to have a nonlinear effect on \( CT_{min} \) (Figure 2a and Figure S8A), the effective degrees of freedom (edf = 1) of the partial effect of latitude suggested a largely linear effect after accounting for altitude (\( F = 40.4, p < .001 \)). Altitude had a more complex effect (edf = 1.262, \( F = 24.0, p < .001 \)), but with a strong altitude outlier in Crested Butte (Table 1), this result should be taken with some caution. Moreover, the overall adjusted R-squared was low at only 0.09. For CCRT, only latitude had a significant effect (edf = 1, \( F = 17.2, p < .001 \); Figure 2b). Taken together, \( CT_{min} \) was lower for populations at higher latitudes and as well as for populations at higher altitudes (Figure 2a and Figure S8A). As one would expect, \( CT_{min} \) showed on average lower values (Figure 2a and Figure S8A), and CCRTs were
shorter (Figure 2b and Figure S8B), at higher latitudes, meaning that more northern and higher altitude populations show higher cold tolerance.

We performed a principal component analysis (PCA) of the WorldClim climate data for a total of 24 *D. montana* populations, from which we had collected samples and where climate data were available. We wanted to examine whether the populations chosen for the cold tolerance tests and genome sequencing were representative of the range of environmental variation experienced by the species. These analyses identified four principal components (PCs) that together explained about 98% of the variation (Figure 3a and Figure S9). The first two PCs separated the populations roughly by a measure of distance from the coast (PC1) and then by latitude and altitude (PC2). PC1 explained ~54% of the variation (Figure 3a and b) and loaded heavily on climate and biological variables associated with precipitation and temperature such as “mean temperature of coldest
cases) make reliable conclusions difficult (see Figure S10). Genomic data were collected. Loadings on PC1 were highly comparable, and although correlations with latitude and altitude did not differ for latitude both in magnitude and direction (rho = 0.6, p = 0.02) and latitude (rho = −0.59, p = .003) correlated with PC2 (see also Figure 3c). Importantly, these patterns were fairly robust when performing PCA using only the six populations for which environmental variations across the six populations for PC1, they covered only ~50% of the range for PC2 (Figure 3a). This suggests that environmental variation across the six populations selected for sequencing reflects that experienced by all populations for PC1. Therefore, the relationship between environmental variables and genetic differentiation in the samples selected for pool-seq is likely to reflect true patterns across populations of *D. montana*, though power might be somewhat reduced for PC2. For all subsequent analyses, we used the results of the PCA using all 24 populations, and to examine the association between climate and phenotype, we compared these across the six common genes are novel to the analysis of PC2 as expected (mean ± SE: 3.6 ± 0.06) by randomly resampling (without replacement) a number of genes equivalent to that associated with top SNPs for each BayScEnv analysis from the *D. montana* annotation, then computing the four-way intersection. This large observed overlap therefore probably reflects a genuine overlap among the underlying top SNPs (data not shown). Some (147, ~13%) of these common genes are novel to *D. montana* (i.e., not annotated in *D. virilis* or in other *Drosophila* spp.) and therefore have no annotation, but 955 have an identifiable *D. virilis* ortholog (Table S4). The second largest set are those genes unique to the analysis of PC2 as an environmental covariate (Figure S15). We performed functional enrichment analyses with *D. montana* annotation, then computing the four-way intersection. This large observed overlap therefore probably reflects a genuine overlap among the underlying top SNPs (data not shown). Some (147, ~13%) of these common genes are novel to *D. montana* (i.e., not annotated in *D. virilis* or in other *Drosophila* spp.) and therefore have no annotation, but 955 have an identifiable *D. virilis* ortholog (Table S4). The second largest set are those genes unique to the analysis of PC2 as an environmental covariate (Figure S15). We performed functional enrichment analyses with *D. montana* keeping all annotation clusters with an enrichment score >1.3 (corresponding to an average corrected p-value of .05). This revealed several common categories of genes associated with the climatic variables and population phenotypes (Table S5). For example, terms associated with membrane and transmembrane structures, immunoglobulins, HAD hydrolase and nucleotide binding were enriched in most of the variables (Table S5). Interestingly, there were also several gene ontology categories that were only enriched in one of the variables, such as glycoside and ATPase hydrolase in *CCRT* and ion channels and transport, as well as metal binding in PC1 (Table S5). *Gowinda* analyses revealed significant enrichment of GO terms (after accounting for variation in gene lengths and correcting for multiple testing) only for genes near SNPs associated with variation in CCRT across populations. Interestingly, the term "carbohydrate derivative binding" was identified among the most enriched terms (Table S6), which agrees closely with some terms identified using *DAVID* for the same variable (Table S5).

### 3.2 Genomic analyses

The number of SNPs with a significant (q-value <0.05) association between population-based *F*~ST~, the two cold tolerance measures (*CCRT* and CTmin), and the two PCs of the bioclimatic data varied from 312 (chromosome 3, *CCRT*) to 2480 (chromosome 4, PC2) across the chromosomes (Figure 4). Using PC1 as an environmental variable with *BAYESCENV* gave a total of 2976 and 1528 SNPs with a q-value <0.05 on the autosomes and on the X chromosome, respectively. Interestingly, the distribution across the chromosomes was not random. Using the distribution of all SNPs to obtain expected counts, there was a significant deviation from expectation ($X^2 = 2906.4, \text{d.f.} = 4, p < .001$). There were many more SNPs than expected on chromosome 4 (1432 vs. 954) and on the X chromosome (1528 vs. 526; Figure 4). Results were similar for PC2 with 6607 and 1861 SNPs with a q-value <0.05 on the autosomes and X chromosomes, respectively. Again, there was a significant deviation from the expected distribution of SNPs across the chromosomes ($X^2 = 1681.9, \text{d.f.} = 4, p < .001$) with an overrepresentation on the fourth (2480 vs. 1794) and the X chromosomes (1861 vs. 989; Figure 4).

We also used average CTmin values per population in similar analyses and found a total of 2668 and 1272 SNPs with a q-value <0.05 on the autosomes and X chromosomes, respectively. The pattern of significant deviations from expected distributions ($X^2 = 2526.2, \text{d.f.} = 4, p < .001$) was also due to an excess on the fourth (1383 vs. 835) and the X chromosomes (1272 vs. 460; Figure 4). Similar results were found for *CCRT* with a total of 2240 and 1228 SNPs with a q-value <0.05, respectively. Once again, there was a significant deviation from the expected distribution of SNPs ($X^2 = 2825.6, \text{d.f.} = 3, p < .001$) with and excess on the fourth (1252 vs. 735) and the X chromosomes (1228 vs. 405; Figure 4). Manhattan plots of the distribution of SNPs across chromosomes are given in Figures S11–S14.

To more closely examine the loci implicated in the four BayScEnv analyses, we identified genes within 10 kb of, or containing the candidate SNPs (Table S4). Overall, there is quite a large overlap among these genes with ~39% (1102 in total) of them being shared by all the four analyses (Table S4, Figure S15). This is far more than would be expected (mean ± SE: 3.6 ± 0.06) by randomly resampling (without replacement) a number of genes equivalent to that associated with top SNPs for each BayScEnv analysis from the *D. montana* annotation, then computing the four-way intersection. This large observed overlap therefore probably reflects a genuine overlap among the underlying top SNPs (data not shown). Some (147, ~13%) of these common genes are novel to *D. montana* (i.e., not annotated in *D. virilis* or in other *Drosophila* spp.) and therefore have no annotation, but 955 have an identifiable *D. virilis* ortholog (Table S4). The second largest set are those genes unique to the analysis of PC2 as an environmental covariate (Figure S15). We performed functional enrichment analyses with *DAVID* keeping all annotation clusters with an enrichment score >1.3 (corresponding to an average corrected p-value of .05). This revealed several common categories of genes associated with the climatic variables and population phenotypes (Table S5). For example, terms associated with membrane and transmembrane structures, immunoglobulins, HAD hydrolase and nucleotide binding were enriched in most of the variables (Table S5). Interestingly, there were also several gene ontology categories that were only enriched in one of the variables, such as glycoside and ATPase hydrolase in *CCRT* and ion channels and transport, as well as metal binding in PC1 (Table S5). *Gowinda* analyses revealed significant enrichment of GO terms (after accounting for variation in gene lengths and correcting for multiple testing) only for genes near SNPs associated with variation in CCRT across populations. Interestingly, the term "carbohydrate derivative binding" was identified among the most enriched terms (Table S6), which agrees closely with some terms identified using *DAVID* for the same variable (Table S5).
S5). Similarly, for PC2, “nucleotide binding” was among the top scoring enriched terms in both GOWINDA and DAVID analyses, though this was not significant after controlling for multiple testing in GOWINDA (Table S5, Table S6).

We then compared loci near candidate SNPs with genes implicated in previous studies of climatic adaptation in *D. montana* including gene expression studies of traits connected to diapause and cold-tolerance (Kankare et al., 2010, 2016; Parker et al., 2015, 2016). Additionally, several candidate genes have been identified near the most significantly differentiated SNPs among *D. montana* populations from Oulanka (Finland), and from North American populations in Colorado and Vancouver (Parker et al., 2018). Finally, quasi-natural selection experiments identified several genes within 10 kb of SNPs responding to selection for a shorter CDL for diapause induction (Kauranen et al., 2019). We tested for an overlap between the total set of genes within 10kb of outlier SNPs from all of the BayScEnv runs (*N* = 2694) and the candidate gene sets identified in earlier studies (see Table S7 for the gene sets and studies used). We computed a bootstrap distribution of overlaps by sampling 2694 random genes from the *D. montana* annotation (*N* = 13,683). For each of the gene sets from previous studies this was done 100 times and the distribution compared to the empirical overlap. Results are given in Table 2. In all the cases the empirical overlap was greater than expected by chance with empirical *p*-values <.05 (ranging from <.0001 to .01; Table 2). The only gene that was found in all five of the previous studies used and in the comparison here is called sidestep II (side-II; Table S8). Unfortunately, there is no information available about the biological processes or molecular functions connected to it. Moreover, from 44 other genes that were common to four of our previous studies and this study (Table S8) most (27) have an ortholog in *D. melanogaster*. These genes have molecular functions such as transmembrane signaling or transporter, acetylcholinesterase, ATP binding, protein serine/threonine kinase, carboxylic ester hydrolase, or Rho guanyl-nucleotide exchange factor activity (Thurmond et al., 2019). Many of the genes are also connected to metal ion, nucleic acid or zinc ion binding (Table S8). After identifying information on molecular or biological function and Interpro domains, eventually only five genes remained for which there was no information available (Table S8).

Figure 4: The observed counts of candidate SNPs (i.e., SNPs with a q-value <.05) across chromosomes for each environmental variable (see Section 2). The total number of SNPs, and the proportion of all SNPs, on each chromosome are given below each set of bars. The expected counts on each chromosome, obtained from the proportions of all SNPs across chromosomes, are shown as points aligned with each bar.

![Figure 4](image.png)

Examination of population genetic parameters identified the Crested Butte population as anomalous. The distribution of Tajima’s *D* is centred close to zero in most populations, being slightly more negative in North American populations (Figure S16A). However, Crested Butte is an outlier with a greatly reduced genome-wide Tajima's *D* (Figure S16A). Similarly, diversity (*a*) is also lower in this population than in other populations. There is no overall relationship between latitude and *a* (Spearman’s *rho* = 0.14, *N* = 6, *p* = .8; Figure S16B) but there is a strong correlation between latitude and Tajima’s *D* which is influenced by this population (with Crested Butte: *rho* = .88, *N* = 6, *p* = .03; without Crested Butte: *rho* = 0.8, *N* = 5, *p* = .13). Although Crested Butte occurs at a much higher altitude (>2800 m) than other populations neither Tajima’s *D* nor *a* correlated significantly with altitude (Tajima’s *D*: *rho* = -0.6, *N* = 6, *p* = .24; *a*: *rho* = -0.6, *N* = 6, *p* = .24). Furthermore, *F*$_{ST}$ was similar across all populations and chromosomes with the exception of Crested Butte which remained an outlier with high *F*$_{ST}$ (Figure 5). Bootstrapped 95% confidence intervals are presented to give a guide to statistical significance of differences among populations and confirm that Crested Butte has high *F*$_{ST}$ (Figure 5). Finally, *F*$_{ST}$ was always highest

| Reference               | N genes | N overlaps | Mean (95% CI) resampled N overlaps | Empirical p-value |
|-------------------------|---------|------------|-----------------------------------|-------------------|
| Kankare et al., (2010)  | 14      | 8          | 2.77 (0, 6)                       | .0004             |
| Parker et al., (2015)   | 42      | 13         | 6.28 (2, 11)                      | .001              |
| Kankare et al., (2016)  | 3,929   | 946        | 773.14 (732, 814)                | <.0001            |
| Parker et al., (2016)   | 130     | 27         | 18.72 (11, 26)                   | .01               |
| Parker et al., (2018)   | 2,114   | 629        | 414.7 (382, 448)                 | <.0001            |
| Kauranen et al., (2019) | 1,751   | 402        | 344.64 (314, 375)                | <.0001            |

Table 2: The overlap of the union of genes within 10 kb of SNPs associated with PC1, PC2, CTmin, and CCRT and previous candidate gene sets (see Table S7)

Shown are the citations for the original study, the number of genes identified from the original study, the number of overlapping genes with the candidate gene set from this study, and the mean and 95% CI from 10,000 resampled sets of candidate genes.
FIGURE 5 Mean, SD, and confidence intervals of the $F_{ST}$ values across each SNP on chromosome and population. $F_{ST}$ was summarised for each SNP by computing all pairwise values then deriving population specific $F_{ST}$ values by averaging across all pairwise values containing that population. The 95% confidence intervals (given by the error bars) are derived from 100 bootstrap samples of SNPs on each chromosome.

4 | DISCUSSION

Detecting genomic signatures of climatic adaptation is an important, but challenging, task. Here we use multiple sources of evidence to study ecological adaptation and population divergence in a highly cold tolerant species of *Drosophila, D. montana*. This species is characterised by a wide circumpolar distribution extending to high latitudes both in North America and Europe, and to high altitudes in the southern part of its range in the Rocky Mountains of North America. These habitats impose extreme seasonal and climatic selective pressure. We collected bioclimatic data from 24 populations along a latitudinal gradient of about 2900 km in North America, and six populations from a gradient of 720 km in Finland. We characterised population level cold-tolerance for six populations from these ranges using two methods, critical thermal minimum (CTmin) and chill coma recovery time (CCRT) and show that CTmin were lower and CCRT shorter in higher latitude populations, as one would expect. Thus, northern populations are more cold-tolerant. Finally, we performed pool-seq of these six populations to investigate the association between genomic and environmental differentiation.

The two methods examining cold tolerance gave somewhat different results, as CTmin, but not CCRT, differed significantly between sexes, with females having lower CTmin than males. In an earlier study investigating seasonal changes in *D. montana* CCRTs, only one out of six comparisons showed a significant difference between sexes (Vesala, Salminen, Kostal, et al., 2012; Vesala, Salminen, Laiho, et al., 2012) and Gibert et al., (2001) did not detect sex-specific differences in CCRT in any of 84 *Drosophila* species they studied. However, several studies of *D. melanogaster* have detected shorter CCRT in females than in males, suggesting that females are more cold tolerant than males (Andersen et al., 2015; Bauerfeind et al., 2014; David et al., 1998), perhaps related to their greater body mass (Wilder et al., 2010). Consequently, the extent and adaptive significance of sex-specific differences in CCRT in *Drosophila* remains unclear.

We derived principal components to summarise WorldClim climatic variables using data from all the 24 populations. The first principal component (PC1) separated populations approximately by a measure of “distance inland” and loaded heavily on climate and variables associated with precipitation and temperature. These results follow the geographic distribution of the populations, for example, the population with highest values for PC1 is Ashford, which is on the Pacific coast and receives most rain, but also experiences warm summers and mild winters. Principal component 2 (PC2), loaded heavily on bioclimatic variables associated with latitudinal clinality, which also mapped onto the populations intuitively as those with higher values on PC2 also occurred at higher latitudes. Interestingly, CTmin values were positively correlated with PC1, but not with PC2, while CCRT showed no relationship with either of these components. This suggests that CTmin and CCRT measure at least slightly different biochemical or physiological mechanisms (Findsen et al., 2014; MacMillan et al., 2012), and could hence be correlated with different climatic variables and also show relatively weak correlation to each other (Andersen et al., 2015). Indeed our results found no significant correlation in CTmin and CCRT across populations. It is also interesting to note that high altitude populations have very similar values on PC1 and PC2 to high latitude populations, most probably reflecting the similar climates in these populations. These similarities in climate are probably also underlying the similarities in CTmin between the high altitude population (Crested Butte) and the high latitude populations from Finland.

The Bayesian analysis identified SNPs showing an association of genetic differentiation with climatic and phenotypic variation. The extent to which the loci associated with the phenotypes and adaptations to different climatic conditions are correlated is one of the most important factors in understanding the evolutionary processes occurring in these populations. The results of our study suggest that the loci associated with the phenotypic measures we have quantified are not strongly correlated with each other and are likely to be under the influence of both genetic and environmental factors. Furthermore, the results suggest that the association between genomic and environmental factors is not strong enough to explain the observed differences in cold tolerance between sexes and populations.

The results of our study also suggest that the genomic signatures of climatic adaptation in *Drosophila* are complex and involve a large number of private genes. The fact that we were able to identify a large number of private genes associated with cold tolerance suggests that the genetic basis of cold tolerance is likely to be highly complex and involve a large number of genes. This is consistent with previous studies that have identified a large number of genes associated with cold tolerance in *Drosophila* (Findsen et al., 2014; MacMillan et al., 2012). The results of our study also suggest that the genomic signatures of climatic adaptation in *Drosophila* are complex and involve a large number of private genes. The fact that we were able to identify a large number of private genes associated with cold tolerance suggests that the genetic basis of cold tolerance is likely to be highly complex and involve a large number of genes. This is consistent with previous studies that have identified a large number of genes associated with cold tolerance in *Drosophila* (Findsen et al., 2014; MacMillan et al., 2012).
is an excellent example of how strong ecological selection may be detected in genomic studies. In particular, because the Bayesian methods examining both ecological variables and relevant phenotypes gave significant overlap amongst the associated loci, and that these are further associated with more broad genomic differentiation between populations, gives confidence that we are consistently identifying genes associated with ecological selection.

Analyses of the functional annotation of these genes strengthen our conclusions that climate driven adaptation is important. Regions near SNPs associated with climatic variables were enriched for genes previously identified as candidates related to cold tolerance, diapause and responses to changes in day length in *D. montana* (Kankare et al., 2010, 2016; Parker et al., 2015, 2016). Thus, genetic variation across populations of these flies may be largely shaped by differences in ecological and climatic variation. This ecological specialisation may have also contributed to the divergence of *D. montana* from its relatives. Parker et al., (2018) surveyed the rates of molecular evolution in eleven cold tolerant and noncold tolerant species of *Drosophila*. The genes found to be evolving at faster rates in cold-tolerant species were enriched for many of the same functional categories as in our current study, including for example, membrane and transmembrane proteins and immunoglobulins (Parker et al., 2018).

The fact that many of the same annotation terms are enriched in clusters for all environmental variables in *DAVID* analyses, suggests that similar genes or biochemical pathways are involved in these adaptations. Membrane proteins and lipids are an important determinant of membrane and cuticular permeability at different temperatures, which in turn has an effect on the resistance to desiccation stress in insects (Gibbs, 2002; Stanziano et al., 2015). Importantly, there is evidence for a close link between the desiccation stress response and cold tolerance across species and in *Drosophila* in particular, suggesting an overlap in some of the mechanisms involved (Sinclair et al., 2007). Cold-hardy lines of *D. melanogaster* are known to exhibit elevated lipid metabolism, perhaps in order to allow rapid lipid membrane modification (Williams et al., 2016) in different environmental conditions. Pleckstrin homology (PH) domain was one of the functional clusters found in both the current study and in the species comparison of Parker et al., (2018). This domain is a flexible module of 100–120 amino acids which interacts with a variety of different ligands, composing a protein-protein interaction platform (Schefzek & Welti, 2012). As changes in the membrane lipid biochemistry form an integral part of the cold tolerance response, genes associated with PH may assist in homeoviscous adaptation namely, alteration of membrane phospholipid composition to maintain fluidity at low temperatures (Sinensky, 1974). Interestingly, the only gene found in all five of our previous studies of cold tolerance in *D. montana* was side-step II (side-II). Unfortunately, there is no information available for the biological processes or molecular functions associated with this gene, but side-II has protein features including immunoglobulin and immunoglobulin-like domain superfamily (Thurmond et al., 2019) and could be involved in immunological processes during cold response of the flies (Parker et al., 2021). Insects are known to produce a diverse range of antimicrobial peptides and proteins as part of their immune activity against viruses, bacteria, fungi and parasites (Mylonakis et al., 2016) and hence immune responses could be part of a general stress response in cold tolerance (Ferguson et al., 2016; Sinclair et al., 2013). We also find confirmation of some enriched GO terms in *GOWINDA* analyses that control for variation in gene length, although overlap is not substantial. In summary, our results indicate that some of the same biochemical processes that are targeted by selection on larger evolutionary scales (i.e., across species), are also involved in local adaptation for different populations within a species, providing a rare bridge between the processes of population differentiation and speciation.

At the chromosomal level, we found an overrepresentation of loci associated with ecological selection on chromosomes X (which corresponds to chromosome 2L in *D. melanogaster*) and 4. It is well-known that X chromosomes can generally evolve quickly due to selection on semi-recessive advantageous loci in the hemizygous sex, and smaller effective population size (Charlesworth et al., 1987) and are often most divergent between closely related species (Abbott et al., 2017; Ellegren et al., 2012). However, there is no obvious reason to expect faster divergence in chromosome 4. *D. montana* populations have considerable chromosomal polymorphism (Morales-Hojas et al., 2007; Patterson & Stone, 1952; Stone et al., 1960) and American populations have been classified into several chromosomal forms on the basis of their geographical origin, size and chromosome structure (Throckmorton, 1982). Moreover, both the X chromosome and chromosome 4 of *D. montana* are known to harbour polymorphic inversions (Morales-Hojas et al., 2007; Stone et al., 1960) and such inversions have often been found to vary clinally in other systems (Chen et al., 2012; Kapun et al., 2016; Kolaczkowski et al., 2011). Recent genomic data confirms that there are several inversions in both chromosomes X and 4, but we do not yet have any detailed information on their distributions within populations (Poikela et al., unpublished data). The reduced recombination within inversions can independently capture advantageous alleles under selection (Kirkpatrick & Barton, 2006) and divergence is often greater in chromosomes carrying inverted regions and noncollinear regions (Lohse et al., 2015) and such regions may diverge more quickly during speciation with gene flow. Indeed, here we found that the X chromosome and chromosome 4 always have the highest levels of *F*ST across all populations, as expected if the ecological selection on loci on these chromosomes influenced overall patterns of genomic diversion, perhaps due to hitchhiking.

Finally, our findings include intriguing result regarding the Crested Butte population which shows reduced genetic variability and a substantial reduction of Tajima’s D relative to the other populations. This population occurs at a very high altitude (>2800 m) and also shows reproductive incompatibilities with other populations (Jennings et al., 2014). It may have also been bottlenecked during its adaptation to this high altitude, so population expansion or selected sweeps could be prevalent within this population (Garlovsky et al., 2020). Whether ecological specialisation is associated with the spread of incompatibilities is an intriguing possibility.
5 | CONCLUSION

Identifying the genetic variation that underlies population divergence and ultimately speciation remains a challenge. Detecting associations between genetic and environmental differentiation at loci across populations has become a popular approach. Here, we apply Bayesian methods to detect such loci across populations of Drosophila montana, which is an extremely cold-tolerant Drosophila species where we expect strong ecological selection. We identify many genes that are associated with both climate variables and population-level cold-tolerance phenotypes. These genes overlap with candidate genes from other studies of variation in cold-tolerance in D. montana and were also overrepresented on chromosomes X and 4 which may be associated with inversion polymorphisms known to be present in these chromosomes. Our study thus provides a clear example of how strong ecological selection can be detected in genome studies, using Bayesian methods to detect local adaptation in combination with studies of ecologically important variables and phenotypes.

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AUTHOR CONTRIBUTIONS

Anneli Hoikkala, Michael G. Ritchie, and Maaria Kankare conceived the study. Venera Tyukmaeva performed the phenotyping experiments. R. Axel W. Wiberg performed all analyses. R. Axel W. Wiberg, Maaria Kankare and Michael G. Ritchie led the writing of the manuscript. All authors contributed to the final version of the manuscript.

DATA AVAILABILITY STATEMENT

Raw reads have been deposited with NCBI (https://www.ncbi.nlm.nih.gov) under the BioProject accession PRJN588720.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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