Selective effects of a short transient environmental fluctuation on a natural population

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Running head

Fluctuating selection in C. riparius

Keywords

Genotype-phenotype map, fluctuating selection, selective tracking

Abstract

Natural populations experience continuous and often transient changes of environmental conditions. These in turn may result in fluctuating selection pressures leading to variable demographic and evolutionary population responses. Rapid adaptation as short-term response to a sudden environmental change has in several cases been attributed to polygenic traits, but the underlying

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genomic dynamics and architecture are poorly understood. In this study, we took advantage of a
natural experiment in an insect population of the non-biting midge *Chironomus riparius* by
monitoring genome-wide allele frequencies before and after a cold snap event. Whole genome
pooled sequencing of time series samples revealed ten selected haplotypes carrying ancient
polymorphisms, partially with signatures of balancing selection. By constantly cold exposing
genetically variable individuals in the laboratory, we could demonstrate with whole genome
resequencing i) among the survivors, the same alleles rose in frequency as in the wild and ii) that the
identified variants additively predicted fitness (survival time) of its bearers. Finally, by simultaneously
sequencing the genome and the transcriptome of cold exposed individuals we could tentatively link
some of the selected SNPs to the *cis*- and *trans*-regulation of genes and pathways known to be
involved in cold response of insects, like *Cytochrome P450* and fatty acid metabolism. Altogether, our
results shed light on the strength and speed of selection in natural populations and the genomic
architecture of its underlying polygenic trait. Population genomic time series data thus appear as
promising tool for measuring the selective tracking of fluctuating selection in natural populations.

5680 words
Introduction

Adaptation in natural populations occurs when selection acts on variable phenotypic traits with a heritable basis. There is a general agreement that selection in the wild is intense (Hoekstra et al., 2001; Kingsolver et al., 2001). It is also variable in space and time (Bell, 2010; Price, Grant, Gibbs, & Boag, 1984; Siepielski, DiBattista, Evans, & Carlson, 2011), even though there is some debate whether changes in the direction of selection are frequent or not (Kingsolver, Diamond, Siepielski, & Carlson, 2012; Kingsolver & Pfennig, 2007; Siepielski, DiBattista, & Carlson, 2009). Recent theoretical (Messer & Petrov, 2013) and empirical work (Bitter, Kapsenberg, Gattuso, & Pfister, 2019) has shown that selection in natural populations can lead to rapid adaptation, in particular of polygenic traits (Barghi, Hermisson, & Schlötterer, 2020; Jain & Stephan, 2017). If the rate of environmental change is not too fast and the population characteristics allow for effective selection, adaptation from standing genetic variation to track moving phenotypic optima is theoretically possible nearly in real-time (Matuszewski, Hermisson, & Kopp, 2015). It is therefore possible that at least some organisms, for example multivoltine species with large population sizes, adaptively track their fluctuating environment (Bell, 2010).

This theoretical basis, however, has currently little empirical support from natural populations. While there are on the one hand examples of selective tracking of the fluctuating environment for phenotypic traits (de Villemereuil et al., 2020; Grant & Grant, 1989; Marrot, Garant, & Charmantier, 2017) and on the other demonstrations of rapid selectively driven changes on the molecular level (Margres et al., 2017; Yang et al., 2016; Zong, Li, & Liu, 2021), we are not aware of studies bringing together the observed temporal fitness differences among different phenotypes with the underlying molecular variants in natural populations.

In this study, we took advantage of a natural experiment to tackle the above question. We studied the genomic response of natural population of a non-biting midge to a short-term weather event, in this case a cold snap. This promised the opportunity to study the selective effects of a defined
transient event as is typical for the selective regime of fluctuating environments (Bell, 2010). Non-biting midges of the Chironomid family are widely distributed aquatic insects and have a crucial role in freshwater benthic ecosystems serving as a basis of benthic food webs (Horváth, Móra, Bernáth, & Kriska, 2011; Oppold et al., 2016; Pfenninger & Nowak, 2008). *Chironomus riparius* (Meigen, 1803) is a multivoltine species with up to 15 generations per year in Europe (Oppold et al., 2016). Therefore, the different generations are subjected to widely varying environmental conditions. Accordingly, extensive research on temperature and photoperiod has shown that several traits can and do adapt locally (Waldvogel et al., 2018), and temporally among seasons (Doria, Caliendo, Gerber, & Pfenninger, 2022; Foucault, Wieser, Waldvogel, Feldmeyer, & Pfenninger, 2018). But also other factors are known to act as selection pressures on this species (e.g. organic load, Kraak *et al.* (2000), conductivity, (Pfenninger & Nowak 2008), nitrogen, (Nemec *et al.* 2012), temperature, (Nemec *et al.* 2013) and anthropogenic substances, (Nowak *et al.* 2009). The high effective and demographic population size (> 1,000,000, Waldvogel *et al.* (2018)) and the very high number of offspring per breeding pair (400-800) allows for rapid adaptation (Pfenninger & Foucault, 2020). Since genomic resources and parameters are available (Schmidt *et al.* 2020; Oppold & Pfenninger 2017) and the species is amenable for evolutionary experiments in the laboratory (Foucault, Wieser, Waldvogel, & Pfenninger, 2019), the species is increasingly becoming a model for molecular ecology and the emerging field of evolutionary ecotoxicology (Doria, Hannappel, & Pfenninger, 2022; Doria, Waldvogel, & Pfenninger, 2021).

In this study we focussed on three research questions. Does normal, transient environmental variation like a cold snap trigger measurable molecular selection in a natural population? Are the putatively selected SNP-loci linked to longer survival also under experimental cold exposure conditions? And finally, can we link the identified variants to lower level phenotypic changes, i.e. gene expression differences?

**Material and Methods**
Temporal sampling of natural population

In the course of routine sampling for another project (Pfenninger & Foucault, 2022), we sampled larvae of the species Chironomus riparius on Feb. 15 2018 with a sieve at a single site situated in a small river (Hasselbach, Hessen, Germany 50.167562°N, 9.083542°E) following the protocol of Foucault et al. (2019b). The sampling site is located close to a wastewater treatment plant (Abwasserverband Freigericht) that continuously monitors physical and chemical water parameters, which they generously provided. A few days after the sampling, the air temperature in the region fell substantially below zero for a couple of days, which eventually drove the water temperatures at the sampling site from the long-term average of 9-10°C during this time of the year down to about 5°C for 2 consecutive days (Figure 1a). We seized the opportunity to obtain another sample of 80 individuals from the same site. Please note that no reproduction takes place in this species at temperatures below ~12-14°C and thus the same generation was sampled. A third sample from the same site was obtained in September 2018, about 6-7 generations later (Oppold et al., 2016). The taxonomic identity of the larvae was ascertained by DNA-barcoding of a mitochondrial (COI) and a nuclear locus (L44). Eighty thus identified C. riparius were pooled and subjected to Pool-sequencing (see below).

Population genomic analyses

DNA was extracted for the three pools from the field using the Quiagen blood and tissue extraction kit on pooled samples of 80 larval head capsules, respectively. Integrity and quality of extracted DNA was controlled using electrophoresis, and the DNA concentration for each samples measured with a Qubit fluorimeter (Invitrogen).

Whole genome pool-sequencing was carried out on an Illumina MiSeq with 250bp paired end reads. Reads were trimmed using the wrapper tool Autotrim (Waldvogel et al. 2018) that integrates Trimmomatic (Bolger et al. 2014) for trimming and FastQC (Andrews 2010) for quality control. The trimmed reads were then mapped on the latest C. riparius reference genome (Schmidt et al. 2020).
using the BWA mem algorithm (Li & Durbin 2009). Low quality reads were subsequently filtered and
SNPs were initially called using Samtools (Li et al. 2009). The pipelines PoPoolation1 v.1.2.2 and
PoPoolation2 v.1.201 (Kofler et al. 2011a; Kofler et al. 2011b) were used to call SNPs and remove
indels. Allele frequencies for all SNPs with coverage between 15x and 70x were estimated with the R
library poolSeq (Taus et al. 2017).

Selected SNP loci were identified by their allele-frequency change (AFC) larger than expected by
sampling variance. Neutral simulations were used to compute false discovery rate q-values < 0.001
with parameters (number of SNPs, starting allele frequencies matching the ancestral population,
sequence coverage, number of generations) matching those of the respective samples. To be
conservative, we calculated the drift for one generational passage. As effective population size, we
used 15,000, which constitutes a very conservative estimate as well (see Waldvogel et al. (2018)).
The resulting -log10 p value includes an assessment of the sampling depth at the respective locus. All
calculations and simulations were performed with the R-library poolSeq (Taus et al. 2017).

To infer the selection regime acting in the long-term at the identified loci and linked sites, we
calculated Tajima’s D as a summary statistics of the site frequency spectrum (Hohenlohe, Phillips, &
Cresko, 2010). As this statistics is insensitive to short-term changes (Hohenlohe et al., 2010), only a
single pool was used for the analysis. We used Popoolation1 to calculate Tajima’s D for all non-
overlapping 1 kb windows in the genome. We then compared Tajima’s D from the windows
harbouring the candidate loci with the distribution of this statistics in all other windows (Schreiber &
Pfenninger, 2021). The window size was chosen based on the short average LD (< 150 bp) in this
species (Pfenninger & Foucault, 2022).

The long-term behaviour of the candidate loci was investigated using data from Pfenninger &
Foucault (2022). The mean allele frequency for each locus was determined for seven time points and
the deviation from this mean was plotted for each time point. In addition, we determined the
average pairwise correlation of these loci over time and compared it with that of the same number
of randomly selected, uncoupled SNPs to see whether the candidate loci tend evolve in concert over longer time periods as well.

Experimental confirmation

To verify the association of the SNPs to survival under cold stress, we exposed 160 4th instar larvae from a genetically variable laboratory population initially gained from the same population (Pfenninger & Foucault, 2022) to 4°C until they died or survived for at least 28 days. The individuals were kept separately in 2 cm well plates with at least 1 cm water column in a normal fridge. We checked daily whether they were still alive by touching them to see if they still moved. Dead larvae or larvae still alive on the 28th day were individually transferred to tubes filled with 70% alcohol and the day of their death recorded. Of these we chose 30 individuals which died early and 30 individuals which died late or even survived until the end of the 28th days for individual whole genome resequencing (see below).

To link genotype to gene expression, we were not able to use the dead individuals from the survival experiment mentioned above. We therefore performed a corresponding short-term experiment exposing another set of 54 4th instar larvae from the same laboratory population to 4°C, this time for three days only in order to guarantee for survival. After these three days, 36 living individuals were cut into three pieces on a -80°C cool pad. We cut two segments from a mid-body segment from each larva for subsequent DNA isolation and resequencing, and the rest of the individual was stored at -80°C for later RNA-isolation. We chose the two mid-body segments for DNA extraction, because these segments are most redundant with other such segments in terms of specialised organs or tissues, thus preventing bias or omission in RNA analysis as far as possible.

DNA/RNA isolation and sequencing

In total, DNA was isolated from 96 individuals (60 from the long-term, and 36 from the short-term 4°C exposure experiments) using the Qiagen® blood&tissue kit. RNA was extracted using the Quick-
RNA Miniprep kit (Zymo Research) from 36 single individuals. Library preparation and 150bp paired-end sequencing was conducted on a NovaSeq platform at Novogene.

Identification of individual genotypes

Quality trimming and mapping of reads was conducted similar to the approach outlined above. Genotypes at the SNP positions identified in the PoolSeq approach were called with bcftools v.1.10.2 (Li, 2011). The genotypes were cross-checked manually for a random sub-sample of individuals with IGV viewer v.2.8.2 (Thorvaldsdóttir, Robinson, & Mesirov, 2013). We calculated the mean number of potentially adaptive alleles (i.e. those that rose in frequency in the natural population) per variable locus (MNAA) for each resequenced individual as quantitative measure of the multi-locus genotype at the respective loci, thereby assuming an additive genotype-to-phenotype relationship (Sella & Barton, 2019).

Association of survival times with genotypes

Under the assumption that alleles identified to rise in frequency during the cold snap indeed conferred collectively a fitness advantage (i.e. polygenic trait of longer survival) to its bearers, we expected that individuals carrying more of these alleles should also tend to survive longer in the experiment. We used a Bayesian implementation of a Pearson correlation analysis (Bååth, 2014) to test for a correlation between MNAA and survival times. Instead of testing the biologically implausible assumption of no association between variables, the Bayesian approach evaluates support for a positive or negative correlation in the data and provides high density intervals, corresponding to confidence limits, for the Pearson correlation parameter estimate (Bååth, 2014).

RNA-Seq analysis of cold-exposed individuals and co-expression networks

Three samples were of bad quality and thus not sequenced. For the other 33 samples, we obtained between 28 to 69 Mio reads per individual, of which ~0.1% of bp were trimmed and 100% of the reads remained after trimming (Supplemental file 2, Table 1). Adapters were trimmed and quality checked with TrimGalore (Krueger, 2016). HiSat2 v.2.1.0 (Kim, Paggi, Park, Bennett, & Salzberg, 2019) was used
to map the reads to the *C. riparius* genome (Schmidt et al. 2020). The counts table (Supplemental file 2, Table 2) was created with HTSeq (Anders, Pyl, & Huber, 2015). To prevent spurious results due to low read counts, we removed genes with less than 10 reads in at least four samples, and samples C4 and C6 due to missing allele frequency information. The differential gene expression analysis was conducted using DESeq2 (Love, Huber, & Anders, 2014) with the mean number of potentially adaptive alleles per variable locus (MNAA) as continuous variable. We tested for association of genotypes at the identified variable SNP positions and normalised gene expression for genes within +/- 200 kb on the same scaffold with the Bayesian correlation test described above.

To identify networks of co-expressed genes (modules), we constructed a weighted gene co-expression network analysis using the R package WGCNA (Langfelder & Horvath, 2008), based on the genes that had passed the quality filtering step for the expression analysis (N = 8,264). Gene counts were normalized using the `varianceStabilizingTransformation` function from DESeq2 (Love et al., 2014). Following the WGCNA guidelines, we picked a soft-thresholding power of 6 for adjacency calculation. To associate modules to the MNAA polygenic score, we first calculated the modules’ eigengene using the `moduleEigengenes` function and tested for module trait correlation using the `corPvalueStudent` function. To obtain up-to-date annotations, we ran a local blastp (Altschul, Gish, Miller, Myers, & Lipman, 2008) of the *C. riparius* proteins versus the non-redundant protein database (version January 2022). We ran Interproscan v.5.53-87.0 (Jones et al., 2014) locally to obtain GO information using the *C. riparius* predicted proteome. The GO enrichment analysis on genes within the significant module was performed with the R package TopGO (Alexa & Rahnenführer, 2016), using the ‘parentchild’ algorithm and the Fishers exact test for significance.

**Results**

**Cold snap and Sampling**

From the 21st of Feb. 2018, the air temperature at the sampling site dropped for 10 consecutive days below zero, with a minimum daily average of -8°C on the 27th of February 2018. The water
temperature, usually fluctuating around 10°C in winter, started to fall slowly as well. At the end of this period, the water temperature dropped steeply to 5°C for two consecutive days. We sampled the population six days after the water temperatures were back to about 10°C (Figure 1A). Sampling the required number of larvae after the cold snap took considerably more time than before.

**Large allele frequency changes within a single generation**

The allele frequency changes of 19 SNPs between before and after the cold snap could not be explained by sampling variance (Figure 1B). These SNPs were therefore considered as candidates for selection (Table 1). Some of these SNPs occurred on the same scaffold in close spatial proximity to other such SNPs (within 40-750 bp, Table 1). As resequencing data showed, the rising alleles at these SNPs were linked to haplotypes (data not shown). We considered the regions with several SNPs therefore as a single locus and the SNP with the largest AFC was used as marker SNP for these linked haplotypes. Taking this into account, ten loci, each on a different scaffold of the reference genome and thus most likely physically unlinked (Pfenninger & Foucault, 2022), were potentially affected by selection. We refer to these loci by their scaffold numbers hereafter (e.g. scaffold227).

The starting frequencies of the rising alleles at these candidate loci before the cold snap were highly variable, spanning the range from undetectably rare (scaffold424 and scaffold674) to the majority allele (scaffold85, Figure 1C). All candidate alleles rose in frequency by at least about 0.5 (0.467-0.781, Table 1, Figure 1C). In September 2018, the allele frequencies of all but one locus (scaffold227) dropped back towards the level they had before the cold snap event (Figure 1C, Table 3).

The effect of the cold snap on allele frequency spectra varied between candidate loci. Comparing Tajima’s D in the 1 kb windows encompassing the selected regions with the distribution of all 1kb windows showed that four selected regions had a Tajima’s D in the upper 5% quantile (scaffold227, scaffold256, scaffold547, scaffold673), indicating balancing selection. One value (scaffold694) was in the lower 5% tail, suggesting a recent selective sweep. The remaining values were inconspicuous (four) or not calculable (one, Figure 1D).
Validation experiment

Of the 160 larvae constantly exposed to 4°C, the first larvae died after 15 days. Mortality on day 21 was extraordinarily high (53 individuals, 34%). After 28 days, 17 (11%) larvae were still alive. This was a much higher mortality than the usually observed 10-25% from hatching to eclosion in experimental settings at normal temperatures (Foucault et al., 2019). For seven individuals, the dying day could not be clearly determined. The distribution of survival over time can be found in Supplemental Figure 1.

Genotyping of experimental individuals

For 59 of the 60 randomly selected individuals from the experiment, resequencing was successful. The individuals could be genotyped on average at 9.02 out of the 10 candidate loci. Two loci were fixed for the rising allele in the experimental sample (Table 3) and were thus not further considered.

In the 59 individuals used in the experiment, all candidate alleles had a considerably higher start frequency than in the natural population before the cold snap (Table 3). Nevertheless, among the survivors at day 25, the frequency of all candidate alleles rose between 0.04 and 0.22 in the course of the experiment with moderate (60.6%) to very high (98.4%) posterior probability (Figure 2A). The mean number of potentially adaptive alleles per variable locus (MNAA) locus ranged between 0.50 and 1.38 among the individuals in the experiment with a mean of 0.98 (s.d. 0.18). Individuals with genotypes containing more potentially adaptive alleles (MNAA) survived the cold stress conditions longer. The association between these variables was moderately strong (most likely estimate of $r = 0.33$, 95% high density interval between 0.08 and 0.55, Figure 2B). The correlation of MNAA with the length of survival was positive with almost certainty (posterior probability 99.3%) and also significantly different from zero in a frequentist approach ($p < 0.01$).

Genotype associations with local and global gene expression data

For 31 individuals with genotype information, we obtained gene expression data for 11,386 of the 13,449 annotated genes (85%). Only two loci showed all possible genotypes at the candidate loci in the individuals used for transcription analysis (scaffold 227 and scaffold 694). On their scaffolds, the
genotypes were strongly associated with the expression levels of one (scaffold 227, Figure 3A), or respectively two (scaffold 694, Figure 3B) genes. On scaffold 227, the selected SNP was in an exon of the gene to which it was associated via the genotype specific expression levels. Specifically, the candidate allele was associated to a higher expression rate. The gene (scaffold227_gene0.177) is annotated as Cytochrome P450, family 6 (CYP6). The two genes with expression levels strongly associated to genotypes on scaffold 694 were both roughly 100 kb away from the selected SNP. The first gene, scaffold694_gene0.209 codes for protein (RFT1) that is essential for protein N-glycosylation and also here the candidate allele was associated with an increase of transcription. In contrast, for the second gene scaffold694_gene0.209, a transmembrane receptor protein tyrosine phosphatase (DEP1), a lower transcription level was associated with the rising candidate allele.

In total, we found the expression of 28 genes to be significantly associated with the MNAA. Nine out of these (32%) belong to genes of (larval) cuticule proteins, endocuticule proteins or endochitinase. One of the 20 distinct co-expressed modules identified, module-cyan (containing 169 genes Supplemental file 2, Table 3), covaried substantially with MNAA (Figure 4A, r = 0.41, p=0.03, Supplemental file 2, Table 4). A GO-enrichment analysis revealed that gene functions related to fatty acid metabolism were overrepresented in this module (Figure 4b, Supplemental file 2, Table 3). The cold snap candidate loci themselves were not part of module cyan.

Discussion

Natural experiment

In this study, we took advantage of a natural experiment. Following a sampling routine, we coincidently sampled a population pool from a natural C. riparius population in late winter just before a cold snap (Pfenninger & Foucault 2020). Such cold snaps at this time of the year are not the rule in Germany, but also not uncommon. The drop of temperature was marked, but not extreme. Likewise, the duration of the snap was relatively short, at least with regard to the temperature drop in the water. It was therefore an event with the potential to leave a selective mark, but it was not an
extreme weather event, let alone a catastrophe. This promised the opportunity to study the selective
effects of a defined transient event as is typical for the selective regime of fluctuating environments
(Bell, 2010). To infer allele frequency changes potentially driven by selection, we therefore sampled
another pool from the same site directly after the cold snap.

*C. riparius* does not reproduce at temperatures below 10°C and larval development is nearly stalled.
We can therefore rule out selection based on differential reproductive success over the time-span of
the experiment (Reznick, 2016). Any potential selection must have occurred in the same generation
by differential mortality of different genotypes. The substantially increased time to collect sufficient
specimen second sampling despite digging deeper indeed indicated that the cold snap most likely
resulted in significant mortality in the population. As chironomids tend to dwell as deep as the
interplay between their haemoglobin concentration and oxygen-availability in the environment
allows (Panis, Goddeeris, & Verheyen, 1996), it was unlikely that the differential catch-rate was e.g.
due to deeper burrowing larvae. Without being able to precisely quantify the demographic decline,
this indicates a much smaller population size after the cold snap, as experience has shown that the
duration of sampling for a predefined number of individuals at a known occurrence site depends on
the population density.

At the same time this meant that genetic drift, defined as sampling variance among generations
(Wright, 1948), could not have played a role here, since there was no generational passage involved.
We have nevertheless included one generation of drift in the calculation of the selection threshold.
This threshold is therefore particularly conservative which increases our certainty that the observed
allele frequency change beyond this threshold is not due to random processes (Barghi et al., 2019;
Hohenlohe et al., 2010). On the other hand, we have certainly missed smaller, yet selection driven
allele frequency changes.

*Signs of selection in the natural population*
In total, ten regions in the genome showed signs of selection in the data. Given our rather conservative threshold, which required a large change in allele frequency (~0.5) for detection, we assume that many loci with less pronounced changes remained undetected. The candidate haplotypes were very short, often a single SNP, which indicated that they are relatively ancient polymorphisms long since segregating in the population (Nordborg & Tavaré, 2002) and thus separated by recombination from the background in which they arose. Analysis of Tajima’s D for the 1kb windows the selected haplotypes resided in showed that nine out of ten had a positive D, four even in the upper 5% quantile. This indicated that the respective polymorphisms could be regularly under differential, balancing selection (Fijarczyk & Babik, 2015). Given that the presumed selection pressure was a seasonal event, a more or less regularly fluctuating environment with opposite selection pressures in winter and summer appeared plausible. Alternatively, the return to initial frequencies may be due to costs associated with antagonistic pleiotropy (Marden, Langford, Robertson, & Fescemyer, 2021). Both can lead to a long-term maintenance of the polymorphism under biologically plausible scenarios (Wittmann, Bergland, Feldman, Schmidt, & Petrov, 2017).

Recent works suggested that balancing selection could be more widespread than previously thought (Gloss & Whiteman, 2016).

The view that seasonal fitness is related to the different alleles at the identified loci was confirmed by the observation that allele frequencies at the candidate loci, with one exception (scaffold 227), returned to near their original frequencies a few generations later. This remarkable correlation between the temporal allele frequency trajectories suggested that the same selection pressure with changing signs among seasons was acting. Seasonally selected polymorphisms with correlated allele frequency trajectories were also observed in natural populations of another dipteran species, Drosophila melanogaster (Bergland, Behrman, O’Brien, Schmidt, & Petrov, 2014; Croze et al., 2017; but see Buffalo & Coop, 2020). But also the selection-driven beak variability of Galapagos finches in response to different weather conditions in different years, taking into account the different generation times, are a classic example of very rapid evolutionary adaptations to a variable
environment (Boag & Grant, 1981). Interestingly, we found a strong negative correlation between the start frequency of the selected SNPs and the absolute deviation from neutrality as measured by Tajima’s D (Supplemental Figure 2). Theory predicts balanced allele frequencies for overdominance (i.e. by heterozygote advantage (Slatkin & Muirhead, 1999)). However, we are not aware of theoretical predictions for expected allele frequencies due to balancing selection by temporally changing selection pressures.

Validation experiment

We observed a substantial variation in survival time in the validation experiment. Compared to the field observations, it took quite a long time (15 days) until the first larvae started to die. In the field, the temperature dropped for two days only and this was obviously long enough to trigger a substantial mortality. This discrepancy could have several, mutually not exclusive explanations.

First, the lab population had already quite high allele frequencies at the loci in question. If these loci were indeed responsible for the longer survival, the lab population could have been a priori better protected against the cold exposure. This shift in allele frequency relative to the natural population might be due to random drift in the relatively large but nevertheless demographically necessarily restricted lab population. However, the high allele frequencies could also be a tribute to the practice of storing egg ropes at 4°C for a few days prior to initial population set-up and experiments to synchronise their development (Foucault et al. 2019). This could have involuntarily preselected the lab population.

Second, the larvae in the experiment came from normal development at benign temperatures (~20°C) and were well-fed before they were exposed to 4°C. The larvae in the field likely hatched in autumn and had passed already several months at about 10°C before the cold snap set in. This resulted in substantial “treatment” differences. Initially, the experimental midges experienced a veritable cold-shock, while the natural population was comparatively slowly cooled down only a few degrees. In Drosophila, it was shown that the rate of temperature change may well have an impact
on the organismal reaction to cold exposure (Gerken, Eller-Smith, & Morgan, 2018). However, the cold shock did not cause any mortality in the experimental population, so that the effect of long-term exposure to low temperatures was the likely major cause for mortality in both experimental and natural population. Probably more importantly, the level of internal resources the two groups could draw upon were likely very different. The laboratory population could draw on their fat reserves, while the natural population likely had nearly exhausted this resource by the end of the winter and thus died faster when exposed to increased cold stress. Lastly, it is likely that the lab experiment did not cover all selection factors that were acting in the field. Given the simplified experimental environment, it is almost certain that the set of selection factors acting on the larvae in the field was different from those in the experiment (Pfenninger & Foucault, 2020).

Despite the reduced complexity of the validation experiment, it confirmed nevertheless the hypothesis that the candidate alleles identified in the natural population conferred longer survival during the experimental cold exposure. The same selection pressure triggered an overall increase in frequency of the same alleles in the experiment by differential mortality as was observed in nature. This is strong evidence that the candidate alleles indeed played a role in the selection process.

The positive correlation between a straightforward polygenic score (mean number of adaptive alleles per locus involved) and the survival time strongly indicated a relatively simple relation: the more of these alleles are present in an individual, the better are the chances of its longer survival under cold stress. Given the likely involvement of more, but yet unidentified loci, nonlinear interactions among loci and non-quantified environmental components (Sella & Barton, 2019), the degree of determination found here appeared quite substantial. Additional evidence that these loci are co-selected by the same selection pressures came from a strong temporal covariance in allele frequencies, also over longer time scales.

Single locus and multi locus genotype associations with gene transcription data
None of the putatively selected SNPs was within the coding region of an annotated gene. We therefore expected phenotypic effects rather due to changes in the transcription regulation of spatially more or less proximate genes than in structural protein changes.

In an attempt to link the identified loci with basal phenotypic aspects, we sequenced a set of non-lethally cold-exposed individuals for both the genome and the transcriptome. We found transcripts for a substantial proportion of the annotated genes. This is similar to results found in *Drosophila* (Brown et al., 2014). With a what could be called “inversed eQTL approach” (Gilad, Rifkin, & Pritchard, 2008; Majewski & Pastinen, 2011), we explored spatially proximate and thus putative cis-interactions between identified selected sites and gene expression levels. By analysing only loci shown to be involved in phenotypic variation and restricting the spatial extent of the search to a plausible range (Schoenfelder & Fraser, 2019), we retained sufficient statistical power even with our relatively limited sample size. Since only two of the identified SNP loci (scaffold227 and scaffold694) showed all possible three genotypes in the sample for which both genotype and transcription data was available, the search for associated cis-regulated genes was necessarily restricted to these.

The gene associated to the selected site on scaffold 227 was identified as Cytochrome P450, family 6, a gene-family that is characteristic for insects (Lewis, Watson, & Lake, 1998). This gene was already several times implicated in the reaction to cold stress (Huang et al., 2017; Lv et al., 2020; Zhang et al., 2015; Zhou, Shan, Tan, Zhang, & Pang, 2019) with an increased expression level under cold conditions. It is therefore plausible that an allele associated with an inherently increased expression level as observed here is under positive selection under cold stress conditions. While the selected marker SNP on scaffold 227 was spatially closely linked to the gene with the associated transcription regulation, the genes most credibly associated to the SNP genotype on scaffold 694 were both roughly 100 kb up-, respectively downstream. Both genes were also not the closest neighbouring genes, but the 4th, respectively 5th transcribed gene up- respectively downstream on the same scaffold. These observations are compatible with recent models of gene-regulation by long range interactions (Schoenfelder & Fraser, 2019) that were also observed in insects (Dorsett, 1999).
gene situated upstream was identified as RFT1 homolog. This protein of the endoplasmatic reticulum membrane appears to be necessary for glycolipid translocation and normal protein N-glycosylation, but its exact function is unknown (Gottier et al., 2017). The associated gene downstream was most similar to a transmembrane receptor protein tyrosine phosphatase DEP1. The selected allele was associated to the downregulation of the respective transcripts. We could not find any studies that have previously linked this gene to cold stress.

Using the polygenic score as a proxy for survival time on the individual level was associated to 28 differentially expressed genes. One third of these (nine) belong to (larval) cuticule proteins, endocuticle proteins or endochitinase, suggesting a role for these genes in differential survival. Moreover, the polygenic score as a proxy for survival time on the individual level revealed a moderately strong correlation (r = 0.4) to a co-expression module. The co-expression module was statistically enriched for genes involved in fatty acid metabolism. The fatty acid metabolism is known to be crucial for the overwintering of insects (Sinclair & Marshall, 2018; Storey & Storey, 2013; Toprak, Hegedus, Doğan, & Güney, 2020), i.e. under cold stress e.g. by providing the necessary energy storage resources or keeping membranes subtle by changing their fatty acid composition (Overgaard, Sørensen, Petersen, Loeschcke, & Holmstrup, 2005). There are many examples showing that unsaturated fatty acids increase under cold temperatures in insects (reviewed in Clark and Worland 2008; Teets and Denlinger 2013). Moreover, fat content was one amongst other phenotypes linked to exposure to a new temperature in Drosophila, which also appears to be a polygenic trait (Barghi et al., 2019).

The association suggested that at least some of the identified loci may be directly or indirectly involved in trans-regulation of the transcription of these genes. A potential candidate for such a trans-regulation was one of the spatially closest genes to a selected site; a histone acetyltransferase p300 on scaffold673. These genes are high level switches that regulate broad scale gene transcription pattern via chromatin remodelling (Tropberger et al., 2013). Amongst many other biological functions, it has been shown that the gene is responsible for the differentiation of fat cells (Gesta,
Tseng, & Kahn, 2007). The transcription level of this gene was significantly associated with the observed genotypes (Supplemental Figure 3), suggesting that a trans-regulated altered transcription of this gene could provide a link between the multilocus genotype and the fatty acid metabolism genes.

This finding allowed us to hypothesise that the identified selected SNPs could, amongst others, contribute to the observed variation in survival time via particular physiological trait(s). Individuals with a high polygenic score for the loci involved could e.g. accumulate more fat reserves, which increased survival time under prolonged cold conditions. This clearly beneficial fitness effect during the winter season could be reversed in summer, when the accumulation of then unnecessary fat reserves deviates resources from reproduction or increases attractivity for predators. Such a more or less regularly fluctuating selection regime on a polygenic trait could well conform to the theoretical preconditions necessary to maintain the respective polymorphisms over longer periods (Wittmann et al., 2017). Whether there is really a link between the extent of the individual fat reserves and the observed survival time under cold stress conditions, however, remains to be tested, just like a causal relation between the respective multilocus genotype and the regulation of the fatty acid metabolism. However, many more intermediate level phenotypic traits, e.g. changes in cuticule composition likely contributed to survival under constant cold exposure.

Conclusions

In this study we could show that normal, short term environmental variability can lead to measurable natural selection on a polygenic trait in a natural population. Time series population genomic analyses from field samples obviously have the power to pick up such transient signals, even if they consist rather of moderate to strong changes in allele frequencies than in fixation of loci. The observed high temporal correlation of the allele-frequency changes of the loci involved holds the promise that changes in the selective regime of natural populations may be identified from population genomic time series data.
Acknowledgements

We thank Andreas Wieser for help with sampling and the Abwasserverband Freigericht for generously providing the environmental Hasselbach data. The work was partially funded by DFG (grant PF390/8-1).

References


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Data Accessibility and Benefit-Sharing

Data accessibility: All sequencing data is publicly available on ENA

PoolSeq data: (project ERP115516, samples ERS4040036-ERS4040041).

Individual Resequencing data: (project PRJEB56138)

Transcriptome data: (project PRJEB56138)

Benefits Generated: Benefits from this research accrue from the research, sharing of our data and results on public databases as described above.

Author Contributions

MP, AMW and BF conceptualised the study, QF carried out the field-work, MP and AMW performed experiments, MP, QF and BF analysed the data, MP drafted the manuscript with contributions from all coauthors.
**Figures**

Figure 1. Field data. A) Air and water temperature curves at the sampling site with sampling dates (grey bars). B) Manhattan plot of the genome-wide SNPs FDR-corrected $-\log_{10}p$ values, contrasting the population pools sampled before and after the cold snap. The horizontal line shows the inferred threshold. C) Allele frequency trajectories at potentially selected loci before and after the cold snap and 6-7 generations later in September 2018. D) Left: Violin-plot of the distribution of Tajima’s $D$ for all 177,185 1 kb windows in the genome. The mean is indicated by a blue horizontal line, the grey area indicates the 90% quantile around the mean. The dashed lines mark the beginning of the upper, respectively lower 5% quantile. Right: Tajima’s $D$ for the 1 kb windows harbouring the candidate SNPs. For one window (scaff396) Tajima’s $D$ could not be computed. E) Long term allele frequency trajectories of the candidate loci. F) Comparison of the mean pairwise correlation coefficient of the candidate loci (left) and random SNPs (right).
Figure 2. Experimental data. A) Temporal course of allele frequency changes at polymorphic candidate loci and survival of individuals during the experiment. Shown in blue are the frequency trajectories of the selected alleles in the natural population. The values to the right show the median Bayesian estimate of increase and the associated posterior probability (pp) of this value being larger than zero for the respective alleles. B) Bayesian estimate of correlation between the mean number of potentially adaptive alleles per locus and the survival time in the experiment.
Figure 3. Associations of genotype at selected sites with gene expression levels of genes within +/- 200 kb on the same scaffold. The position of the selected SNP is indicated with a red arrow. Annotated genes are indicated as blue bars, whose length is proportional to the length of the gene. For genes with expression data available, the genotype-specific transcription level is given in a plot. Within plots, the bars represent the standard deviation range of gene expression variation for the possible genotypes, from left to right: homozygous falling allele, heterozygous, homozygous rising allele. Above the panels, the Bayesian statistic for association are given: $r =$ coefficient of association, HDI = 95% high density interval, $pp =$ posterior probability for the association coefficient being larger or smaller than zero, respectively. Highlighted in red are genes with a HDI not comprising zero and $pp > 95%$. A) Scaffold 227 with selected SNP at position 185,498 with a upregulation associated with the rising allele at gene 0.177. The SNP is positioned in an exon of the gene. B) Scaffold 694 with selected SNP at position 289,921. One gene (0.209), situated about 100 kb from the SNP showed a strong upregulation associated with the rising allele. Another gene (0.262), more than 100 kb away showed a strong downregulation.
Figure 4. Association between Mean Number of Adaptive Alleles (MNAA) and gene co-expression modules. A) Correlation of MNAA with inferred gene co-expression modules. The modules carry arbitrary colour names. In the right column, the Pearson correlation coefficient between individual transcription levels and the polygenic score and its probability of being identical to zero (in brackets) for the respective module is given. For immediate visual recognition, the correlation coefficient was also translated into a heatmap from green (-1) over white (0) to red (+1). B) Word-cloud of significantly over-represented GO-terms in the cyan gene co-expression module. The font size is proportional to the number of genes. GO-terms with relation to fatty acid metabolism are underlined.
### Tables

Table 1. Outlier SNPs in field data.

<table>
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<th>rising allele</th>
<th>start allele freq.</th>
<th>end allele freq.</th>
<th>AFC</th>
<th>-log10p</th>
<th>p fdr-corrected</th>
<th>Tajima’s D 1k</th>
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</thead>
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<td>616771</td>
<td>T</td>
<td>C</td>
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<td>G/alt</td>
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<td>G/alt</td>
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<td>0.591</td>
<td>0.591</td>
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<td>T/alt</td>
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<td>T/ref</td>
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<td>T/alt</td>
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Table 2. Nearest genes to potentially selected loci.

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<th>distance to closest gene</th>
<th>gene start</th>
<th>gene end</th>
<th>annotation</th>
<th>Annotation spatially most proximate gene</th>
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<td>5044</td>
<td>621815</td>
<td>629417</td>
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<td>120</td>
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<td>973122</td>
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<td>6810</td>
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<td>196989</td>
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<td>6914</td>
<td>8455</td>
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<td>24716</td>
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<td>26768</td>
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<td>uncharacterized protein LOC109541342 isoform X2</td>
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<td>297539</td>
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<td>protein dachsous</td>
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Table 3. Allele frequencies in the experiment. AF = allele frequency.

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<th>rising allele</th>
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<th>AF Mar18</th>
<th>AF Sep18</th>
<th>AF in exp pop</th>
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<tbody>
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<td>0.386</td>
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<td>0.800</td>
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<td>A/alt</td>
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<td>0.857</td>
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