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Evolutionary origins of genomic adaptations in an invasive copepod

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The ability of populations to expand their geographical ranges, whether as invaders, agricultural strains or climate migrants, is currently one of the most serious global problems. However, fundamental mechanisms remain poorly understood regarding factors that enable certain populations, such as biological invaders, to rapidly transition to novel habitats. According to one hypothesis, environmental fluctuations in the native range could promote successful invasions by imposing balancing selection on key traits and maintaining the genetic variation that enables rapid adaptation in novel habitats. Here we test the genomic predictions of this hypothesis by performing whole-genome sequencing of multiple independent invasive freshwater and native saline populations of the copepod *Eurytemora affinis* complex. We found that invasive populations have repeatedly responded to selection through the parallel use of the same single-nucleotide polymorphisms and genomic loci, to a much greater degree than expected. These same loci were enriched for signatures of long-term balancing selection in the native ranges, with 15-47% of loci exhibiting significant signatures of balancing selection. The strong association between parallel evolution in the invaded range and balancing selection in the native range supports the hypothesis that fluctuating habitats can promote invasive success and that balancing selection might serve as a widespread and important mechanism that enables rapid adaptation in nature.

nvasive species pose one of the greatest threats to biodiversity, ecosystem integrity, agriculture, fisheries and public health, with economic costs amounting to hundreds of billions of dollars per year worldwide^{1,2}. Global climate change is projected to increase the number and impact of invaders in an unprecedented and complex manner³⁻⁸, requiring a comprehensive understanding of the mechanisms that facilitate successful biological invasions⁹⁻¹². A longstanding debate has focused on the precise factors that generate successful invaders, given that an exceedingly small proportion of introduced species are able to establish in new habitats and then become invasive¹³. Numerous hypotheses have been proposed and tested, including the roles of propagule pressure, transport opportunity, habitat matching, fecundity and population size. However, these hypotheses have not found consistent empirical support across taxonomic groups and invasion events, offering limited powers of prediction¹⁴⁻¹⁸.

Lee and Gelembiuk¹⁹ proposed an evolutionary mechanism that could promote the emergence of invasive populations and hypothesized that the selection regime in the native range acts as a crucial factor that affects invasive success¹⁹. They observed that invasive populations tend to originate from habitats marked by disturbance or temporally varying conditions^{19,20}. Consequently, they hypothesized that many invasive populations have originated from native populations undergoing balancing selection, resulting from fluctuating environmental conditions. This mechanism would tend to operate in organisms with short generation times, relative to the period of environmental fluctuations, such that different alleles would be favoured by selection in different generations¹⁹. Such a selection regime could maintain standing genetic variation in the native range and provide the genetic substrate upon which positive selection could act during invasions^{10,15,17,21-24}. However, this hypothesis had not previously been tested empirically.

Balancing selection is a form of natural selection that favours more than one allele at a locus, and its ability to maintain standing genetic variation has remained a contested topic in evolutionary biology²⁴⁻³⁰. In particular, the conditions under which temporally fluctuating selection can maintain polymorphisms through time have been thought to be fairly restricted³¹⁻³⁵. Moreover, little empirical evidence exists regarding the extent to which adaptation to novel habitats could be facilitated by balancing selection in the native range^{24,36-38}. In theory, balanced genetic variants might have a higher probability than neutral variation of contributing to adaptation to novel habitats, as balancing selection can maintain variants at relatively high frequencies and increase their fixation probabilities under new selection pressures^{24,39-42}. Recently, several studies of adaptation from standing genetic variation have found signatures of directional selection acting on alleles that segregate at intermediate frequencies, suggesting the presence of some form of balancing selection^{38,42-45}. However, whether alleles under directional selection in colonizing or invasive populations could arise from those maintained under balancing selection in their native ranges remains largely untested.

Moreover, balancing selection in the native range could increase the chances of selection acting on the same loci during replicated invasion events. When closely related populations are independently exposed to the same novel selection pressure, adaptation could potentially proceed from shared standing variation⁴⁶. The prevalence of parallel selection acting on shared standing variation depends on the divergence time between populations and the factors that influence the retention of ancestral variation⁴⁷. Fluctuating selection due to shared environmental forces in the native range could result in multiple native range populations harbouring the same ancestral genetic variation. The presence of these shared balanced variants would increase the probability of genetic parallelism during invasions, thereby enhancing the predictability of evolutionary responses to environmental change⁴⁸.

The common estuarine and saltmarsh copepod *E. affinis* complex provides an excellent model system in which to explore these

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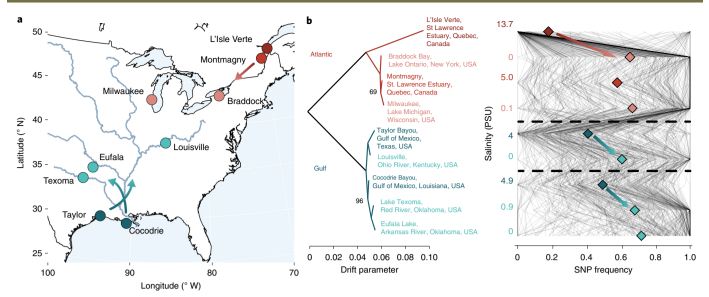


Fig. 1 Population genomic signatures of parallel freshwater invasions. a, Map of sampling locations of *E. affinis* complex populations in North America. Invasive freshwater populations are represented by light-coloured circles and native saline populations are represented by dark-coloured circles. Red circles represent populations from the Atlantic clade and green circles represent populations from the Gulf clade⁵⁴. Arrows indicate the inferred directions of independent freshwater invasions. **b**, Parallel frequency shifts associated with repeated saline to freshwater invasions (parallel candidate SNPs, Supplementary Box 1). The population phylogeny was estimated from SNP frequency correlations using TreeMix v.1.13. Invasive freshwater populations are shown as light colours and native saline populations are shown as dark colours. All nodes have bootstrap support of 100%, except those that are shown. SNP frequencies (grey lines) are shown for parallel candidate SNPs (n=347) with both significant signatures of directional selection (BayeScan 3) and association with salinity (BayPass). SNP frequencies are polarized to keep directionality consistent among SNPs. Diamonds represent the mean candidate SNP frequencies of the corresponding population on the phylogeny. The numbers next to population names indicate the salinity at which each population was sampled. Dotted horizontal lines delineate clades with both invasive and native populations, indicating the presence of recent independent invasions. The arrows represent the direction of SNP frequency shifts between saline and freshwater populations.

questions. Within the past approximately 70 years, populations from this species complex have been very successful at invading freshwater habitats, mediated by human activity. Due to increases in shipping and ballast water discharge in recent years, E. affinis complex populations have invaded freshwater habitats multiple times independently, from genetically divergent clades throughout the Northern Hemisphere^{20,49} (Fig. 1). These replicated invasions enable the discovery of genomic signatures that are consistently associated with invasion success. Notably, native range saline populations of the E. affinis complex exhibit several properties that could greatly expand the conditions under which balancing selection can maintain polymorphisms, including seasonal fluctuations in salinity, overlapping generations in the form of diapause egg banks $^{\rm 50-52}$ and beneficial reversal of dominance with respect to salinity tolerance⁵³ (see 'Discussion' and Box 1). Thus, balancing selection might provide a plausible mechanism that facilitates invasions in this system, and potentially many other systems that give rise to invasive populations¹⁹.

Thus, in this study, we took advantage of the independent invasions and rapid parallel physiological evolution in the *E. affinis* complex to test the hypothesis that fluctuating habitats promote invasion success by imposing balancing selection on key traits and associated genomic loci. To test this hypothesis, we (1) explored genomic targets of directional selection during independent saline to freshwater invasions and assessed the prevalence of molecular parallelism across the genome, and (2) examined whether the specific genomic targets of selection in the invasive populations exhibit signatures of balancing selection in the native populations.

To address our hypotheses, we performed whole-genome sequencing of 100 pooled individuals per population for multiple invasive populations and their respective native range populations (Fig. 1a). This evolutionary replication gave us greater power to determine the specific loci that undergo both directional and balancing selection. We illustrate that the processes and mechanisms that underlie freshwater invasions can be highly predictable, both in terms of generating successful invaders in the native range and inducing parallel evolutionary responses in the invaded range. If indeed the same loci that undergo parallel selection in the invaded range are also under balancing selection in the native range, such information could offer unprecedented powers of prediction on which populations have the capacity to invade.

Results

Population structure and history. Our population genomic data support the occurrence of multiple independent invasions from genetically divergent source populations, corroborating previous studies using mtDNA^{49,54} (Fig. 1a). The deep phylogenetic split between the Atlantic and Gulf clades was evident from the small proportion of shared variation present and the structure of the population phylogeny built from the variance-covariance matrix of single-nucleotide polymorphism (SNP) frequencies⁵⁵ (Fig. 1a). In terms of shared variation in the two clades, only 5.52% of the global 6,635,765 biallelic SNPs had a minor allele frequency (MAF) > 0.05 in both clades. In addition, we found much greater SNP differentiation among populations in different clades (mean $F_{CT} = 0.19$) than within clades (mean $F_{SC} = 0.04$) (estimated using BayeScan 3 (also known as BayeScanHierarchical)⁵⁶, see 'Widespread genomic signatures of parallel directional selection during freshwater invasions'). The population phylogeny indicated at least two independent freshwater invasions in the Gulf clade and one invasion in the Atlantic clade. Genome-wide genetic diversity estimates (θ) (Supplementary Table 2) were not significantly lower in the invasive populations (phylogenetic generalized least squares; $\theta_{\text{Watterson}} \sim \text{salin-}$ ity, t = -0.167, d.f. = 9, P = 0.872; θ_{π} ~salinity: t = -0.335, d.f. = 9, P=0.748; see Methods), indicating the lack of population bottlenecks following invasion events.

Box 1 | Factors that could promote balancing selection in native populations of the *E. affinis* complex

Fluctuating environmental conditions Native populations of the *E. affinis* complex experience seasonal fluctuations in salinity ranging from 5 to 40 PSU²⁰. With around six generations per year, the period of salinity fluctuations is greater than generation time, exposing different generations to different selection pressures throughout the course of the year.

Negative genetic correlations In the *E. affinis* complex, negative genetic correlations (consistent with antagonistic pleiotropy) exist between saltwater and freshwater tolerance⁹⁹⁻¹⁰¹. Thus, under seasonally fluctuating salinities, saline and freshwater tolerance would be favoured by selection at different times and different generations (with around six generations per year). Such fluctuating selection could promote the maintenance of genetic variation, provided that this polymorphism can remain protected against strong negative selection¹⁰²⁻¹⁰⁴.

Beneficial reversal of dominance Beneficial reversal of dominance (BRD) with respect to salinity tolerance has been demonstrated in the *E. affinis* complex⁵³. BRD is the phenomenon in which alternate alleles are always dominant in the environment in which they have a higher fitness. Under fluctuating selection, BRD can greatly increase the fitness of heterozygotes and protect polymorphisms, as alleles will have reduced exposure to selection during conditions in which they are not beneficial^{70,102–107}. Currently, BRD has been demonstrated in only the *E. affinis* complex⁵³, although there is evidence that dominance in gene expression may be environmentally dependent in *Drosophila*⁷⁸.

Overlapping generations Populations of the *E. affinis* complex have overlapping generations generated through diapause egg banks⁵⁰⁻⁵², which can protect polymorphisms against fluctuating selection over time^{102,105}.

Phenotypic plasticity Populations of the *E. affinis* complex exhibit plasticity in physiological tolerance and performance associated with salinity changes in the native range¹⁰¹. Plasticity has been predicted to contribute to the ability of balancing selection to maintain polymorphisms through a genomic storage effect^{108,109}.

Widespread genomic signatures of parallel directional selection during freshwater invasions. Genomic signatures of repeated freshwater invasions were enriched for signatures of directional selection both on the same SNPs and on different SNPs in the same genomic windows in both clades (see Supplementary Box 1 for definitions of relevant terms). This pattern of selection on many of the same loci was surprising given the deep phylogenetic split that we estimated between the two distinct clades of the *E. affinis* complex (Fig. 1a). A substantial number of SNPs and genomic windows displayed signatures of selection in only one clade, suggesting some independent genomic routes to freshwater adaptation. However, using both neutral simulations and window randomizations to generate null distributions, we found that the genome-wide signatures of repeated evolution at the same loci were significantly greater than expectation (Supplementary Information section I).

To detect SNPs associated with freshwater invasions, we performed genome-wide scans for directional selection and association with salinity with BayeScan^{56,57} and BayPass⁵⁸, respectively. A significant 'association with salinity' indicates that the shift in allele (SNP) frequency was correlated with changes in salinity, suggesting a functional relationship between the rise of particular alleles and freshwater adaptation. In addition to detecting genomic signatures of freshwater invasions in each clade separately, we sought to detect signatures of directional selection (Supplementary Box 1) found in common in both clades with the goal of increasing power to detect loci truly associated with freshwater adaptation^{46,59,60}. We used two approaches to detect signatures of selection found in common between the two clades given their apparent divergence: (1) we tested for signatures of parallel directional selection and association with salinity on shared SNPs (to uncover parallel candidate SNPs; Supplementary Box 1) and (2) detected 10-kb genomic windows that overlapped between the two separate genome scans for selection in each clade (shared candidate windows; Supplementary Box 1). The first analysis was designed to detect signatures of parallel frequency shifts at exactly the same SNPs, whereas the second analysis was designed to detect small genomic regions that contained shared targets of selection. Detecting shared candidate windows could capture different SNPs under selection that occur at the same loci in different lineages. As a result of our analyses, we obtained four sets of candidate loci (Supplementary Box 1).

To assess the extent of parallel frequency shifts in shared SNPs, we used a hierarchical F-model (BayeScan 3)⁵⁶ to compare statistical support for three selection models and a neutral model for all SNPs with a MAF>0.05 in both clades (n=366,781). The three selection models were constructed to determine whether a SNP had a signature of directional selection in only the Atlantic clade, in only the Gulf clade, or in both clades in parallel. A substantial proportion of SNPs with significant signatures of directional selection showed the highest support for the parallel selection model, rather than selection in either clade alone (parallel, 42.5%; only the Atlantic clade, 19.5%; only the Gulf clade, 38.0%). In terms of the numbers of significant SNPs, 2,970 SNPs displayed signatures of parallel directional selection out of a total of 6,981 SNPs with signatures of directional selection in at least one clade. Of the 2,970 SNPs with signatures of parallel directional selection, 349 SNPs also showed significant association with salinity across all populations. We removed 2 of these 349 parallel candidate SNPs (Supplementary Box 1) from downstream analyses because they showed evidence of potential copy-number variation (Supplementary Information section II). We found that genome-wide signatures of parallelism for shared SNPs were significantly greater than the degree of parallelism expected under genetic drift alone, based on neutral simulations (Supplementary Information section I, Supplementary Table 3). Specifically, our dataset contained 29 times the number of SNPs with signatures of parallel selection and association with salinity than expected under drift alone (Supplementary Information section I).

The analyses of directional selection and association with salinity in each separate clade yielded 679 and 2,092 non-parallel candidate SNPs in the Atlantic and Gulf clades, respectively. The greater number of non-parallel candidate SNPs found in the Gulf clade was likely due to the power gained from the additional population and invasion event sampled in the Gulf clade relative to the Atlantic clade (Fig. 1). Notably, the number of shared candidate windows (n=279; Supplementary Box 1) around significant SNPs was significantly greater than expected using window randomization $(n=120.31\pm0.204, P<0.0001;$ Supplementary Information section I). Thus, the genome-wide signature of selection on the same small genomic regions was 2.3-fold greater than expected by chance given the size of the windows and genome. While the greater number of non-parallel, compared with the number of parallel, candidate SNPs could indicate largely different genomic routes to freshwater adaptation, the greater than expected signatures of selection both at the same SNPs and at different SNPs in the same 10-kb windows suggest that selection has often acted on the same loci across repeated invasion events and that these parallel loci represent important genomic signatures that underlie the adaptation to freshwater habitats.

Candidate loci	Gene names of the ion transporters
Parallel candidate SNPs	NHA, paralogues 3, 4, 5, 7; NKA, subunit α, paralogue 2; NKA, subunit β, paralogue 5; Rh , paralogue 2
Shared candidate windows	NHE, clade X, paralogue c; NHA, paralogue 6; NKA, subunit α , paralogue 2
Non-parallel candidate SNPs, Atlantic clade	NHA, paralogues 1, 2, 4, 6, 7; NKA, subunit α , paralogues 2, 4, 5; NKA, subunit β , paralogue 5; carbonic anhydrase (CA), paralogues 1, 5, 12
Non-parallel candidate SNPs, Gulf clade	Ammonia transporter (<i>AMT</i>), paralogue 3; <i>Rh</i> , paralogue 2; <i>NKA</i> , subunit α , paralogues 2, 6; Na ⁺ ,K ⁺ ,2Cl ⁻ cotransporter (<i>NKCC</i>), paralogues 3, 4
See Box 1 for definitions of the te	rms.

Ion transporter genes are overrepresented as targets of selection. Ion transport and related Gene Ontology (GO) terms were overrepresented in our candidate SNPs (Supplementary Table 4). Among significant GO terms (false-discovery rate (FDR)-adjusted *P*<0.05), 37.5% (3 out of 8) and 55.5% (5 out of 9) were related to ion transport in the set of parallel candidate SNPs and non-parallel candidate SNPs of the Atlantic clade, respectively (Supplementary Table 4). Top GO categories in the parallel set included several ion transporter terms, such as 'sodium ion transport', 'sodium-proton antiporter activity', 'lithium-proton antiporter activity' and 'regulation of intracellular pH'. Ion transport had previously been implicated in freshwater adaptation in the E. affinis complex⁶¹⁻⁶³, and here our GO analysis also implicates ion transport as the dominant physiological function associated with freshwater invasions. Other top GO terms in the set of parallel candidate SNPs included several terms related to gene regulation (for example, 'protein maturation by protein folding', 'regulation of gene expression', 'RNA strand annealing activity', 'protein O-linked glycosylation', 'histone acetyltransferase complex'), energy production (for example, 'mitochondrial membrane'), stress response (for example, 'response to acid chemical'), immune response (for example, 'respiratory burst involved in inflammatory response', 'response to histamine', 'defense response') and metabolism (for example, '4-hydroxyproline metabolic process', 'positive regulation of protein metabolic process', 'negative regulation of gluconeogenesis').

Our candidate SNPs occurred in genomic regions within or proximate to several manually annotated ion transporter genes (Table 1, Supplementary Tables 5–8). Notably, the highest density of parallel candidate SNPs was found on scaffold 68 in a region that contains seven tandem paralogues of the Na⁺/H⁺ antiporter (*NHA*) (Fig. 2). Parallel candidate SNPs were found within or proximate to four paralogues of the Na⁺/H⁺ antiporter (*NHA* 3, 4, 5, 7), the α and β subunits of Na⁺,K⁺-ATPase (*NKA* α 2, *NKA* β 5) and one paralogue of the ammonium transporter Rh protein (*Rh* 2). Non-parallel candidate SNPs were found within or proximate to additional ion transporter genes, including multiple paralogs of *CA*, *NKCC*, *NHE*, *NKA* and *NHA* (Table 1).

SNPs with signatures of directional selection in the invading range are enriched for signatures of balancing selection in the native ranges. Our population genomic data support the hypothesis that a significant proportion of the genetic variation that responded to directional selection during freshwater invasions was maintained in the native populations by balancing selection. If the genetic variants that facilitated freshwater invasions were maintained by selection due to seasonally fluctuating salinity in the native ranges, this signature of long-term balancing selection should be detectable in the native saline populations. To test this prediction, we scanned the genomes of the saline populations for signatures of long-term balancing selection using two recently developed summary statistics, $\beta^{(2)}$ and Non-central Deviation (NCD2)⁶⁴⁻⁶⁶, using all SNPs with a MAF>0.05 in each population. These newer methods have been shown to be considerably more powerful and robust to non-equilibrium demographic histories than traditional tests, such as Tajima's D^{67} . High $\beta^{(2)}$ scores indicate an excess of SNPs at similar frequencies, while low NCD2 scores indicate a build-up of SNPs near a specified intermediate frequency, both of which are potential consequences of long-term balancing selection. Both statistics also increase in significance with a deficit of substitutions relative to an outgroup. We first tested whether parallel and non-parallel candidate SNPs and 10-kb windows around our candidate SNPs were enriched for signatures of balancing selection relative to the whole genome in each native saline population. We then tested whether the SNPs with the strongest signatures of balancing selection in the native ranges were enriched for signatures of directional selection and association with salinity in the invading freshwater populations.

Notably, we found that candidate SNPs and windows with signatures of directional selection and association with salinity were enriched for signatures of long-term balancing selection in all native saline populations. Specifically, parallel candidate SNPs had significantly stronger signatures of balancing selection than the whole genome in four (using $\beta^{(2)}$) or three (using NCD2) of the native populations (Fig. 3, Supplementary Table 9, Extended Data Figs. 1 and 2). Mean $\beta^{(2)}$ scores were up to 3.65-fold higher in parallel candidate SNPs than the genome-wide average. On average, $\beta^{(2)}$ scores for SNPs in 10-kb windows around parallel candidate SNPs were 3.1-fold higher than the genome-wide average, and $\beta^{(2)}$ scores for SNPs in shared candidate windows were 2.35-fold higher than the genome-wide average. This enrichment of balancing selection in windows around parallel candidate SNPs and in shared windows was significant in all four native saline populations (Supplementary Table 9). Although the enrichment was less pronounced when using NCD2, there was a significant reduction in NCD2 scores (indicating stronger signatures of balancing selection) in all four native range populations (Supplementary Table 9). There was an average decrease in NCD2 score of 7.4% in 10-kb windows around parallel candidate SNPs and a decrease in NCD2 score of 4.9% in shared candidate windows.

In a complementary manner, SNPs with the strongest signatures of balancing selection in the native populations (the top 1% of the $\beta^{(2)}$ distributions and bottom 1% of the NCD2 distributions in each saline population) were significantly enriched for signatures of directional selection and association with salinity relative to the whole genome when considering SNPs shared by both clades (Supplementary Table 9). The one exception was the native range population in Baie de L'Isle Verte, where SNPs in the top 1% of the $\beta^{(2)}$ and NCD2 distributions were not enriched for signatures of directional selection, but were enriched for association with salinity (Supplementary Table 9). SNPs in the top 1% of $\beta^{(2)}$ scores and bottom 1% of NCD2 scores were, on average, 1.29 deciban units higher than the whole genome in the BayeScan 3 analysis. This value corresponds to a 1.35-fold BF increase in support for directional selection for these SNPs under balancing selection. Similarly, these SNPs had, on average, 1.34-fold higher support for association with salinity.

Overall, the strongest signatures of balancing selection were found in the parallel candidate SNPs rather than in shared windows or non-parallel candidate SNPs. Non-parallel candidate SNPs were also broadly enriched for signatures of balancing selection, with up to a 2.43-fold mean $\beta^{(2)}$ score increase in non-parallel candidate SNPs relative to the whole genome (Supplementary Table 10). Notably, parallel candidate SNPs showed stronger signatures of balancing selection than non-parallel candidate SNPs in two (using $\beta^{(2)}$)

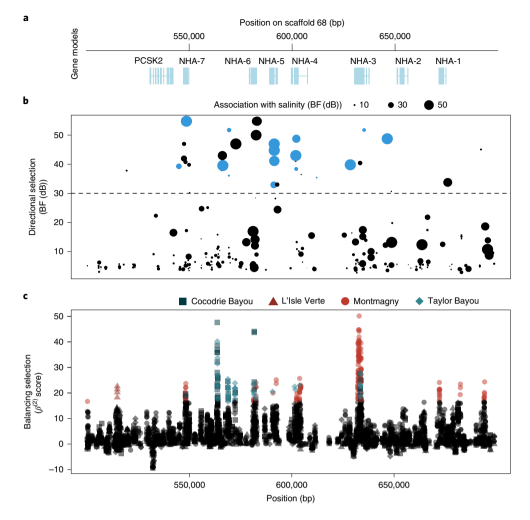


Fig. 2 | Signatures of directional and balancing selection at the *NHA* **cluster in the** *E. affinis* **complex genome. a**, Annotated genes on scaffold 68 of the *E. affinis* complex (Atlantic clade) draft genome⁸³, showing seven tandem paralogues of the *NHA* gene family. **b**, Support for directional selection was estimated with BayeScan 3 by comparing SNP frequencies among the native saline and freshwater invading populations of both the Atlantic and Gulf clades. The horizontal dotted line corresponds to the chosen significance level for directional selection. Blue dots correspond to the SNPs with significant support for parallel directional selection in the BayeScan 3 analysis. Large blue dots (BF>30, deciban units (dB)) above the dotted line are also significantly associated with salinity and are designated as the 'parallel candidate SNPs' (Supplementary Box 1). Black dots above the dotted line correspond to SNPs with support for directional selection in only one clade. The size of the dots corresponds to the level of BF support for association with salinity. **c**, Signatures of balancing selection (based on $\beta^{(2)}$ scores) in the four native saline populations (Fig. 1) from the Atlantic (red triangles and circles) and Gulf (green squares and diamonds) clades. All SNPs in each population for which $\beta^{(2)}$ scores were calculated are shown. Coloured data points represent SNPs in the top 1% of $\beta^{(2)}$ scores in each population.

or three (using NCD2) native range populations (Supplementary Table 11, section A, rows 1-4, 9-12). Similarly, 10-kb windows around parallel candidate SNPs had stronger signatures of balancing selection than 10-kb windows around non-parallel candidate SNPs in all four native range populations (Supplementary Table 11, section A, rows 5-8, 13-16). For example, in the native population in Cocodrie Bayou, $\beta^{(2)}$ scores for parallel candidate SNPs were 4.95-fold higher than non-parallel candidate SNPs and 1.34-fold higher than SNPs in shared windows (Supplementary Table 11, section A, row 3 and section C, row 3). These results suggest that many selected SNPs, comprising both parallel and non-parallel SNPs, were maintained by balancing selection in the native range. However, finding stronger signatures of balancing selection in the set of parallel candidate SNPs than in the set of non-parallel candidate SNPs supports the hypothesis that balancing selection specifically promotes parallel evolution by maintaining shared adaptive variation.

Parallel candidate SNPs exhibited significant signatures of balancing selection in the native range populations to a greater degree than expected using both neutral simulations and whole-genome distributions as null models to calculate significance. In fact, between 15.6% and 47.4% of parallel candidate SNPs fell in the top 5% of simulated balancing selection scores in the native range populations, and between 6.4% and 17.4% fell in the top 5% of the genome-wide distributions (Fig. 3c, Supplementary Table 12). These proportions of parallel candidate SNPs with significant signatures of balancing selection were broadly greater than the genome-wide expectations of around 15% based on the neutral simulation approach and 5% based on the whole-genome outlier approach (Fig. 3c, Supplementary Table 12). Considering the $\beta^{(2)}$ score, approximately 66.0% (using neutral simulations to calculate significance) and 12.5% (using the whole-genome distributions to calculate significance) of 10-kb windows around parallel candidate SNPs exhibited strong signatures of balancing selection in at least one native range population in both clades. These windows with signatures of balancing selection in both clades overlapped with several manually annotated ion transporter genes, including NHA

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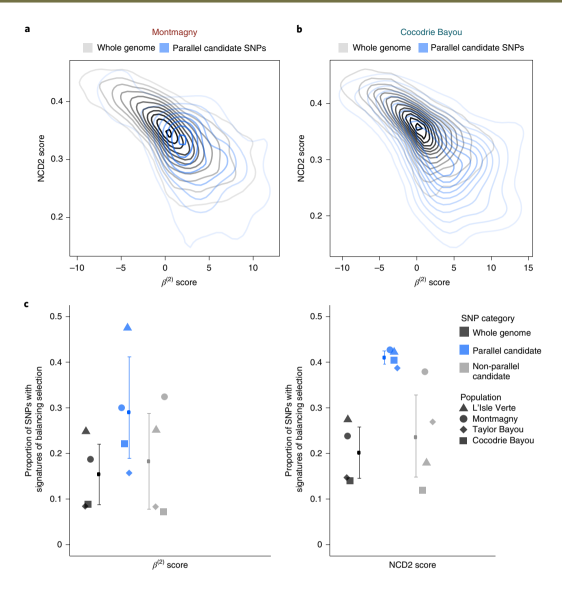


Fig. 3 | Loci with signatures of parallel directional selection and association with salinity are enriched for signatures of long-term balancing selection in the native ranges. Higher $\beta^{(2)}$ scores and lower NCD2 scores indicate stronger signatures of long-term balancing selection. On average, $\beta^{(2)}$ scores are higher and NCD2 scores are lower for parallel candidate SNPs (blue) relative to the rest of the genome (grey). **a**, $\beta^{(2)}$ and NCD2 score density plot for the native, saline Montmagny population (St Lawrence estuary, Atlantic clade). **b**, $\beta^{(2)}$ and NCD2 score density plot for the native, saline Cocodrie Bayou population (Gulf of Mexico, Gulf clade). **c**, Proportion of SNPs with strong signatures of balancing selection based on $\beta^{(2)}$ scores (left) (falling in the top 5% of the simulated null distribution) and NCD2 scores (right) (falling in the bottom 5% of the simulated null distribution). Data points are displayed for each of the four native saline populations (Fig. 1). A greater proportion of parallel candidate SNPs (blue data points; Supplementary Box 1) show signatures of balancing selection, relative to non-parallel candidate SNPs (grey points) and SNPs across the whole genome (black points). Error bars represent 95% confidence limits for the mean values of the four populations.

paralogues 6 and 7, and *NKA* β subunit, paralogue 5. Overall, these results indicate that loci with signatures of long-term balancing selection in the native ranges have a higher probability of responding to directional selection during invasions relative to loci without such signatures.

Discussion

A comprehensive understanding of the evolutionary genetic mechanisms that underlie successful invasions is key to the prediction, prevention and management of current and future biological invasions^{5,9}. In this study, we present evidence for highly repeatable evolutionary genomic mechanisms that underlie rapid adaptation during invasions. Our study provides evidence, at the genomic level, that fluctuating habitats can promote successful invasions of novel habitats by maintaining the genetic variation required for rapid adaptation. We show that a substantial proportion of the genetic variants that underlie freshwater invasions are likely maintained by balancing selection in the native ranges, rather than by mutation-drift balance. In other words, the SNPs and genomic windows with signatures of balancing selection have higher probabilities of responding to directional selection during invasions than loci without signatures of balancing selection. Furthermore, our data provide evidence that balancing selection can enable parallel adaptation to repeatedly deploy the same standing variation preserved in the native range. The marked degree of parallelism at the genetic level appears to be the direct result of balancing selection maintaining a common pool of adaptive genetic variation in the native range populations.

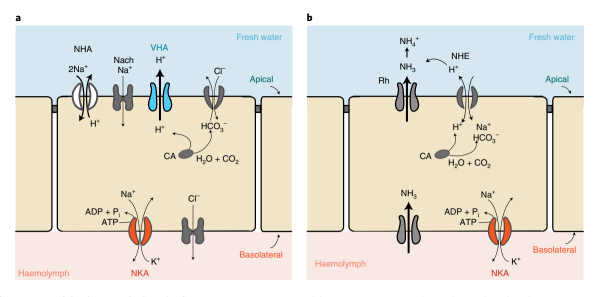


Fig. 4 | Alternative models of ion uptake from freshwater environments. a, Model 1. VHA generates an electrochemical gradient by pumping out protons, to facilitate uptake of Na⁺ through an electrogenic Na⁺ transporter (possibly NHA or Na⁺ channel (Nach)). **b**, Model 2. An ammonia transporter Rh protein exports NH₃ out of the cell and this NH₃ reacts with H⁺ to form NH₄⁺. The reduced extracellular H⁺ concentrations causes NHE to export H⁺ in exchange for Na⁺. These models are not comprehensive for all tissues or taxa and they are not mutually exclusive. Adapted from Lee et al.⁶³ and Dymowska et al.⁹⁸.

Despite the deep phylogenetic distance among the invasive populations in different clades, directional selection often acted on the same loci across invasion events. This parallelism was observed both at different SNPs within the same small genomic windows (possibly representing different SNPs at the same loci) and at exactly the same SNPs in both clades. Although a considerable number of loci had signatures of selection in only one clade, the signatures of selection on shared standing genetic variation was surprising given the large genetic divergences between the native range populations from different clades. Indeed, the detected proportion of candidate SNPs with support for signatures of parallel selection (42.5% when considering shared variation) was higher than several other well-documented cases of parallel evolution68 (for example, 31% of SNPs underlie parallel high-altitude tolerance in humans⁵⁶ and 35% of genomic regions underlie parallel freshwater adaptation in the threespine stickleback⁵⁹). In addition, our analysis found the strongest enrichment of balancing selection among parallel candidate SNPs rather than non-parallel candidate SNPs. Thus, we posit that many of those SNPs had remained segregating in the native populations for so long, making them available for natural selection in multiple populations, because they were maintained by balancing selection from the time preceding the split between the two clades.

According to classical theoretical studies, balancing selection as a result of temporally varying environments has been thought to be restricted in its ability to maintain polymorphisms through time, as the maladapted alleles would be removed from the population by negative selection^{33-35,69}. Even as recent theoretical and empirical studies have suggested that fluctuating selection can maintain variation at many loci70-72, evidence for this phenomenon has remained limited. Notably, native range populations of the E. affinis complex do exhibit several properties that would greatly expand the conditions under which balancing selection could maintain polymorphisms (Box 1). These features, along with the results presented here, make balancing selection a plausible mechanism for maintaining the observed key genetic variation required for adaptation during invasions. While we are not arguing that this mechanism is solely responsible for promoting successful invasions, this study provides strong empirical support that this mechanism is quite possible. Indeed, a large proportion of candidate SNPs did not have

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signatures of balancing selection in the native range, suggesting that neutral and rare variants also play an important role. Nevertheless, the significant enrichment for balancing selection among both parallel and non-parallel candidate loci does strongly suggest that balancing selection is a key mechanism promoting adaptation to novel habitats. Future studies should explore whether seasonal fluctuations in salinity do indeed induce correlated fluctuations in SNP frequencies for candidate loci, producing the genetic signatures of long-term balancing selection for critical traits.

Temporally fluctuating selection is a widespread phenomenon in general^{73,74} and the copepod *E. affinis* complex is not unique among biological invaders in facing environmental heterogeneity in their native ranges¹⁹. For instance, invasive zebra mussel populations have been shown to originate from brackish estuaries marked by heterogeneity, rather than from the more stable ancient lakes75,76. Mechanistically, seasonal changes in dominance can greatly increase the extent of variation maintained by balancing selection, imposed by temporally fluctuating conditions, by protecting maladapted alleles from negative selection^{70,77} (Box 1). While empirical evidence for environmentally dependent dominance has been demonstrated in the E. affinis complex⁵³ and in Drosophila⁷⁸, further work is needed to assess the pervasiveness of beneficial reversal of dominance in nature. Overlapping generations present a potentially prevalent mechanism that protects polymorphisms from negative selection during fluctuating conditions, given that dormancy, seed banks and diapause eggs are found commonly in nature, including in invasive populations⁷⁹. As global climate change promises to modify the extent of spatial and temporal environmental heterogeneity, the ability to detect balancing selection in genomes may become a critical component of predicting future invasions, as well as responses to climate change.

The ability of temporally fluctuating habitats to promote success in introduced populations likely depends on the precise traits under balancing selection in the native habitats and how they relate to new selection pressures. In the present case, ion uptake appears to be a key physiological trait underlying rapid transitions from saline to freshwater habitats^{61,62} (Table 1, Supplementary Table 4). However, fundamental mechanisms of ion uptake in fresh water continue to be debated, particularly the mechanism by which Na⁺ is

transported into cells from very low external ionic conditions (see Fig. 4 for alternative hypotheses). Although our population genomic analyses implicate genes that are possibly involved in both of these models of ion uptake (for example, *NHA*, *NKA* or *Rh*), the strongest and most consistent signal of selection in both the Atlantic and Gulf clades occurred within the *NHA* paralogue cluster (Fig. 2). These results point to a key role of Na⁺/H⁺ antiporter and other ion-transporter paralogues in adaptive ion uptake in fresh water (Table 1, Fig. 4a). Future functional and phylogenetic studies within the Crustacea and across animals would reveal the functions of the ion-transporter paralogues and their general importance for adaptive evolution across salinity boundaries.

In conclusion, we have leveraged deep population genomic sampling of native and invasive populations of the E. affinis complex to uncover both parallel and clade-specific signatures of selection in response to freshwater invasions. We discovered that directional selection in the invasive populations frequently acted on shared standing polymorphisms with signatures of balancing selection in the native ranges. Our results point to a high degree of repeatability in the evolutionary processes that generate successful invaders, from the conditions in the native habitats to the selection response following introductions. Our findings suggest that balanced polymorphisms might often be an integral component of rapid adaptation to novel environments, contributing to a long-standing debate regarding the role of balancing selection in maintaining genetic variation within populations. Given that balancing selection may be more common in nature than previously thought, future studies of adaptation to new environments should consider the role of balancing selection in maintaining critical standing genetic variation upon which selection could act, especially for populations experiencing rapid environmental change.

Methods

Population sampling, raw read processing and SNP calling. The copepod E. affinis constitutes a cryptic species complex, comprising at least six geographically separated and genetically divergent clades distributed across the Northern Hemisphere^{54,80}. Population sampling targeted nine wild populations of the E. affinis complex from two genetically distinct lineages (the Gulf and Atlantic clades)^{49,54} (Fig. 1; Supplementary Table 1). Only the Atlantic clade has been named E. carolleeae⁸¹, but is referred to as the Atlantic clade in this paper for consistency, given that the other clades of the E. affinis complex (for example, the Gulf clade) have not been separately named. In total, five invading freshwater (0-0.9 practical salinity units (PSU), which is approximately equal to parts per thousand salinity) populations and four native saline (>4 PSU) populations were used in this study. From each population, 100 adult individuals were sampled with an approximate 1:1 sex ratio. Individuals were pooled and whole-genome shotgun sequenced (that is, Pool-seq) using the Nextera DNA library preparation kit (Illumina). Libraries were sequenced on an Illumina HiSeq platform at the Institute for Genome Sciences, University of Maryland School of Medicine, generating an average of 179 million 100-bp read pairs per population. Raw reads were trimmed and filtered of adapter sequences, low-complexity sequences and low-quality (Q < 15) bases using BBDuk in the BBTools package (https://jgi.doe.gov/data-and-tools/bbtools/). Processed reads were mapped to the repeat-masked⁸² E. affinis complex (Atlantic clade) draft reference genome⁸³ using BWA-MEM v.0.7.17 (ref.⁸⁴). Paired-end reads that did not align concordantly with BWA-MEM were aligned as single-end reads using NextGenMap v.0.5.5 to aid in the alignment of diverged sequences⁸⁵. The combined read-mapping procedure achieved a mean mapping rate of $95.09 \pm 1.42\%$. Duplicate reads were removed using Picard v.2.18.27 (http://broadinstitute.github. io/picard) and regions around insertions or deletions were realigned using GATK v.3.8 (ref. 86). SAMtools v.1.3.1 was used to convert BAM files into mpileup format after removing low-quality alignments and bases (Q < 20). Sites within 3 bp of an insertion or deletion were removed and the filtered mpileup was converted to sync format using PoPoolation2 (ref. 87). The R package poolfstat v.1.0 (ref. 88) was used to detect bi-allelic SNPs with a global MAF > 0.05, at least four reads were required for a base call, and a minimum of 20 and a maximum of 200 total read counts were required for all populations. In total, 6,635,765 SNPs passed these filters when considering all nine populations. A total of 7,565,621 and 5,323,780 SNPs were called for the Atlantic and Gulf clades, respectively.

Estimating population history. The software TreeMix v.1.13 was used to estimate a bifurcating population tree from bi-allelic SNP frequencies using windows of 1,000 SNPs in size⁵⁵. Only SNPs with a MAF > 0.05 in both clades (that is, shared

variants) were used for this analysis as the model does not consider new mutations. Node support was assessed using 100 bootstrap replicates.

To assess the potential importance of genetic drift during invasions, parameters indicative of effective population size ($\theta = 4N_{d}\mu$) were estimated for each population. PoPoolation⁸⁹ was used to estimate $\theta_{\text{Watterson}}$ and θ_z in 25-kb non-overlapping windows under an infinite sites model. These estimates used the correction for Pool-seq sampling, requiring at least four reads for a base call and a minimum of 20 and maximum of 200 total read counts for a SNP call. Phylogenetic generalized least squares regression was used to test for a relationship between genome-wide θ estimates and salinity. Phylogenetic generalized least squares analyses were performed in the R package phytools v.0.6 (ref. ⁹⁰) with the function pgls.SEy using the phylogeny estimated with TreeMix. Mean θ was regressed against square-root-transformed salinity, accounting for variance in θ for each population using a previously published method⁹¹.

Detecting genomic signatures of parallel directional selection in the invading populations. BayeScan 2 (ref. ⁵⁷) was used to detect signatures of directional

populations. Bayescan 2 (ref. -) was used to detect signatures of directional selection in each clade separately—that is, to identify the non-parallel candidate SNPs (Supplementary Box 1). The hierarchical version of BayeScan (that is, BayeScan 3 or BayeScanHierarchical⁵⁶) was used to detect SNPs that display parallel signatures of directional selection in both the Atlantic and Gulf clades—that is, parallel candidate SNPs (Supplementary Box 1). For the detection of parallel candidate SNPs, the dataset was restricted to SNPs with a minimum MAF of 0.05 in both clades (that is, excluding alleles that are fixed in either clade), such that 366,781 SNPs remained.

In each BayeScan analysis, the Markov chain Monte Carlo (MCMC) chain was run three times independently with 20 pilot runs of 500 iterations each, a burn-in of 2,500 iterations and 1,000 samples collected with a thinning interval of 50 iterations. As parameter estimates were highly consistent across runs, parameter estimates were taken as the median of the three runs to remove potentially spurious estimates from any individual run. The prior odds for selection were set to 0.01 to match the default prior of the BayPass model58 to facilitate a direct comparison of the BF results from the two models. For the BayeScan 3 analysis, the support for a model with selection versus neutrality was taken as the sum of the posterior probabilities of the three selection models (only the Atlantic clade, only the Gulf clade and parallel). These posterior probabilities for selection were then converted to BFs in deciban units (10log₁₀(posterior odds/prior odds)). BF values greater than 30 deciban units were considered decisive evidence for selection, and the model of selection (only the Atlantic clade, only the Gulf clade or parallel) with the highest posterior probability was identified as the best fit model. For the BayeScan 2 analyses, BF values were estimated from the α parameter and converted to deciban units as above. To analyse the large number of SNPs in the independent clade analyses, each dataset was subsampled into sets of around 100.000 random SNPs.

The BayPass v.2.1 package⁵⁸ was used to identify SNPs with frequencies that were significantly associated with salinity. Posterior distributions of the parameters of the Pool-seq version of the BayPass models were estimated for each SNP using the following MCMC procedure: 15 pilot runs of length 500 were run before a burn-in of 2,500 iterations and a sampling of 1,000 MCMC samples with a thinning interval of 25 iterations. To analyse the large number of SNPs in the independent clade analyses, each dataset was subsampled into sets of approximately 100,000 random SNPs. Parameter estimates were taken as the median of the three independent runs. To estimate the expected variancecovariance matrix for SNP frequencies, the posterior distributions of the parameters of the BayPass core model were estimated using the aforementioned MCMC procedure. To estimate the correlation between each SNP and salinity, posterior distributions of the parameters of the auxiliary model were estimated, considering salinity measured at the time of sample collection as the environmental covariate (Supplementary Table 1). As recommended⁵⁸, salinity was scaled so that the mean = 0 and variance = 1. The auxiliary δ parameter estimates the probability that the frequency of a SNP is correlated with a variable of interest. The δ parameter estimates were used to calculate BFs using the default prior distribution58. SNPs displaying both a significant (BF > 30) association with salinity and support for directional selection were considered to be candidate SNPs (Supplementary Box 1) that were putatively linked to targets of selection during freshwater invasions. Parallel candidate SNPs were also examined for aberrant coverage profiles indicating potential mapping artefacts that could result from copy-number variation or mapping biases due to the divergence between the two clades (Supplementary Information section II). Candidate SNPs with such signatures (n = 2) were removed from downstream analyses.

Approximate gene annotations were obtained by assigning SNPs to their closest gene model in the *E. affinis* complex (Atlantic clade) genome using BedTools v.2.28 (ref. ⁹²) and BEDOPS⁹³. Putative ion-transport-related genes were manually annotated with the Web Apollo platform⁹⁴ using a combination of BLAST searches against the NCBI RefSeq and nr databases, i5K Arthropod Genome databases, FlyBase and FleaBase. To determine the specific paralogue clade identities for our candidate genes, phylogenies were constructed of candidate gene families across the Arthropoda as described previously⁸³.

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GO enrichment tests were performed to detect functional groups that were enriched in each of our three categories of significant SNPs (Supplementary Box 1). We assigned GO terms to *E. affinis* complex gene models using significant (E < 0.001) BLAST hits in the Uniprot/Swissprot database. We used Gowinda⁹⁵ to detect GO terms that were significantly enriched in our candidate SNPs, using 100,000 simulations to assess significance. We assigned GO terms to SNPs falling within 10kb of gene models (that is, including SNPs in potential regulatory regions) and counted only one SNP per coding region to account for linkage. As the tests were designed to be applied to SNP datasets, they were not performed on shared candidate windows. *P* values were corrected for multiple tests using the Benjamini–Hochberg procedure⁹⁶.

Detecting genomic signatures of balancing selection in the native range populations. To test whether the SNPs under directional selection in the freshwater invading populations were maintained by balancing selection were performed for each of the four native range populations using the $\beta^{(2)}$ and NCD2 statistics. In contrast to traditional tests, such as Tajima's *D* and the Hudson–Kreitman–Aguade test, these methods explicitly consider the expected allele frequency distribution under long-term balancing selection. High $\beta^{(2)}$ scores indicate an excess of SNPs at similar frequencies^{64,66}, while low NCD2 scores indicate a build-up of SNPs near a specified intermediate frequency⁶². Both statistics also increase in significance with a deficit of substitutions relative to an outgroup.

 $\beta^{(2)}$ and NCD2 scores were calculated for SNPs with a MAF > 0.15 to reduce false positives⁶⁴⁻⁶⁶. This MAF threshold was determined previously⁶⁴, based on simulations suggesting that balanced SNPs were unlikely to achieve equilibrium frequencies <0.15 due to their high probability of drifting out of the population. A MAF filter of 0.05, evaluated for each population separately, was applied to SNPs in the windows used to calculate the scores. SNP frequencies were polarized and substitutions were called using the alternate clade as the outgroup. A window size of 500 bp was used based on the estimated recombination rate of the copepod Tigriopus californicus97. The NCD2 statistic was developed to calculate scores for genomic windows, while $\beta^{(2)}$ calculates scores for individual SNPs. To facilitate comparison between the two statistics, a custom Python script was used to calculate a modified NCD2 statistic for every SNP (rather than genomic window) in the saline population genomes using windows of 500 bp around each SNP and considering a target frequency of 0.5. Although this target frequency was chosen arbitrarily, NCD2 scores have been shown to be robust to the choice of target frequency65

To estimate a null distribution of $\beta^{(2)}$ and NCD2 scores under a model with only genetic drift, we calculated $\beta^{(2)}$ and NCD2 scores for SNPs simulated under the parameters of the neutral variance–covariance matrix (that is, population history). Simulated read counts (that is, SNP frequencies) were generated using the R function simulate.baypass included in the BayPass v.2.1 package using the core BayPass model parameters inferred from the full SNP dataset. The simulated SNPs were placed in the empirical SNP positions along the genome to calculate the null distribution of $\beta^{(2)}$ and NCD2 scores for each native range population. Empirical values were then compared with these null distributions to determine deviation from the null model as a measure of significance. As our simulation approach ultimately assumed that a large proportion of the genome was subject to balancing selection, we also used more conservative empirical cut-offs based on the whole-genome distributions.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

Raw sequence data and aligned reads have been deposited in the NCBI Sequence Read Archive under BioProject ID PRJNA610547.

Code availablility

Custom scripts used throughout this analysis are available online at https://github.com/TheDBStern/NEE2020.

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Author contributions

C.E.L. contributed to the design of the study and data collection. D.B.S. contributed to the conception and execution of the analyses. Both authors contributed to the writing and approval of the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

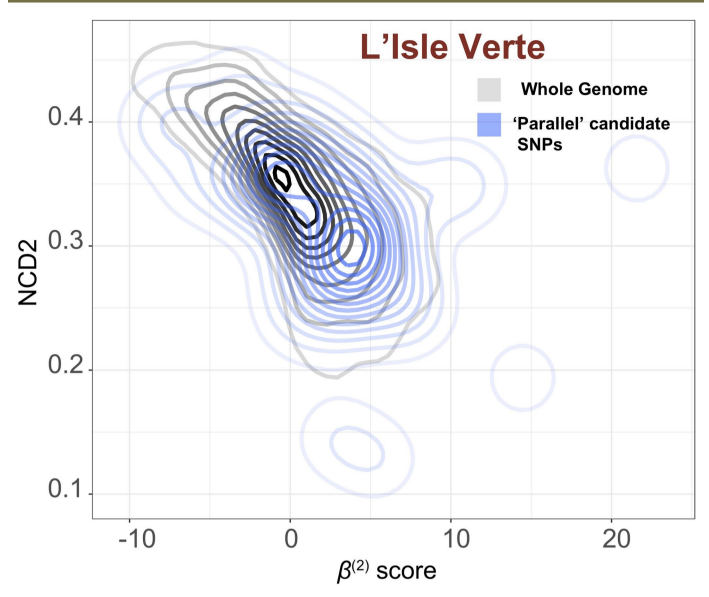
Extended data is available for this paper at https://doi.org/10.1038/s41559-020-1201-y. **Supplementary information** is available for this paper at https://doi.org/10.1038/

s41559-020-1201-y. Correspondence and requests for materials should be addressed to C.E.L.

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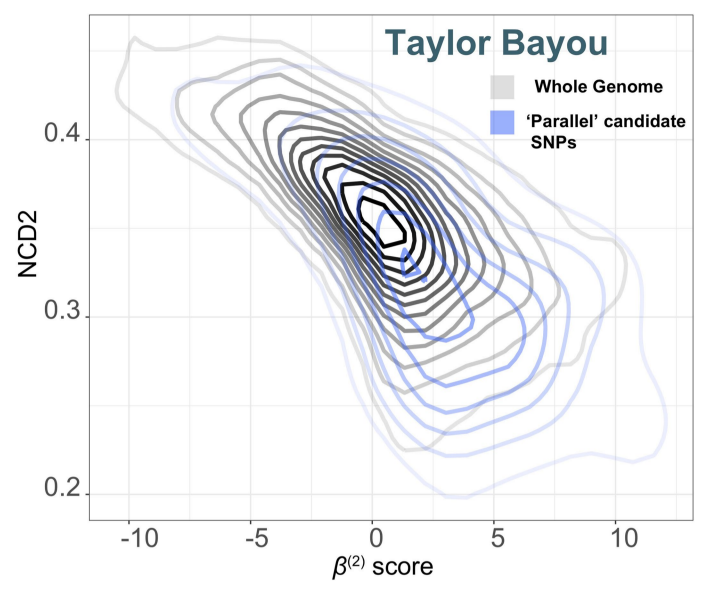
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Extended Data Fig. 1 $\beta^{(2)}$ and NCD2 score density plot for the native, saline Baie de L'Isle Verte population (St. Lawrence drainage, Atlantic clade). Higher $\beta^{(2)}$ scores and lower NCD2 scores signify stronger signatures of long-term balancing selection. $\beta^{(2)}$ scores are higher and NCD2 scores are lower on average for 'parallel' candidate SNPs (blue) relative to the rest of the genome (grey).

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Extended Data Fig. 2 | $\beta^{(2)}$ and NCD2 score density plot for the native, saline Taylor Bayou (Gulf of Mexico, Gulf clade). Higher $\beta^{(2)}$ scores and lower NCD2 scores signify stronger signatures of long-term balancing selection. $\beta^{(2)}$ scores are higher and NCD2 scores are lower on average for 'parallel' candidate SNPs (blue) relative to the rest of the genome (grey).

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		Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.

Software and code

Policy information a	bout <u>availability of computer code</u>
Data collection	No software was used for data collection except for software built into the Illumina HiSeq platform.
Data analysis	BBTools v38.33, BWA-MEM v0.7.17, NextGenMap v0.5.5, Picard v2.18.27, GATK v3.8, SAMtools v1.3.1, PoPoolation v1.2.2, PoPoolation v1.013, poolfstat v.1.0, TreeMix v1.13, phytools v.0.6, BayeScan 2, BayeScan 3, BayPass v2.1, BetaScan2, GenMap, edgeR, BedTools v2.28, BEDOPS, Gowinda, custom code (https://github.com/TheDBStern/poolseq_utils)

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Raw sequence data and aligned reads have been deposited to the NCBI Sequence Read Archive under BioProject ID PRJNA610547.

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Ecological, evolutionary & environmental sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description	This paper describes a re-sequencing study of nine wild populations of copepod Eurytemora affinis complex. 100 individuals were sampled from each population and pooled prior to sequencing. Patterns of genetic diversity within and among populations were analyzed to detect signatures of natural selection.
Research sample	Multiple native and invaded range populations of the copepod Eurytemora affinis complex were sampled. The specific populations were chosen because there were hypothesized to represent multiple independent invasion events from genetically distinct sources, providing evolutionary replication.
Sampling strategy	The number of individuals sampled per population (N=100) and sequencing depth was determined based on recommendations from prior studies (e.g. Schlötterer et al. Nat. Rev. Gen. 2014). The number of populations / localities sampled was to maximize the number of replicated invasion events in this species.
Data collection	DNA sequencing data were collected using the Nextera DNA library preparation kit (Illumina, Inc., San Diego, CA, USA). Libraries were sequenced on an Illumina HiSeq platform at the University of Maryland, School of Medicine, Institute for Genome Sciences. Salinity measurements from each sampling site were collected using a handheld refractometer.
Timing and spatial scale	All tissue and DNA samples were collected within the same year and sequenced at the same time.
Data exclusions	No data were excluded from this analysis.
Reproducibility	All of the scripts needed to reproduce these results are provided in a publicly accessibly Git repository.
Randomization	Animals were allocated into groups based upon sampling location.
Dlinding	Plinding was not relevant to this study, as the relevant data (SNP frequencies) were collected via cognerating machine
Blinding	Blinding was not relevant to this study, as the relevant data (SNP frequencies) were collected via sequencing machine.
Did the study involve field	d work? 🛛 Yes 🗌 No

Field work, collection and transport

Field conditions	Field conditions were typical of the locations and time of year.
Location	Sampling locations are presented in Supplementary Table 1
Access and import/export	The subjects of this study were highly abundant, non-regulated aquatic invertebrates and we did not require permits for collection.
Disturbance	Collections were made using plankton tows, a non-invasive / non-disturbance causing collection method.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

Antibodies

Clinical data

 \boxtimes

 \square

Methods

n/a Involved in the study n/a Involved in the study \square ChIP-seq \boxtimes Eukaryotic cell lines Flow cytometry \boxtimes MRI-based neuroimaging Palaeontology Animals and other organisms Human research participants

Animals and other organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research		
Laboratory animals	No laboratory animals were involved in this study	
Wild animals	Copepods were collected using plankton tows. Plankton was brought back to the lab and sorted. Eurytemora affinis complex were pooled and subjected to DNA extraction. Other plankton properly disposed of.	
Field-collected samples	No manipulative laboratory experiments were performed.	
Ethics oversight	No ethics approval was required because this study used highly abundant, non-regulated invertebrate specimens.	

Note that full information on the approval of the study protocol must also be provided in the manuscript.