# A layover in Europe: reconstructing the invasion route of asexual lineages of a New Zealand snail to North America

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July 16, 2020

#### Abstract

Non-native species are threatening ecosystems and biodiversity worldwide. High genetic variation is thought to be a critical factor for invasion success. The global invasion of a few clonal lineages of the gastropod *Potamopyrgus antipodarum* is thus both puzzling and has the potential to help illuminate why some invasions succeed while others fail. Here, we used SNP markers and a geographically broad sampling scheme (N = 1617) including native New Zealand populations and invasive North American and European populations to provide the first widescale population genetic assessment of the relationships between and amongst native and invasive *P. antipodarum*. We used a combination of traditional and Bayesian molecular analyses to demonstrate that New Zealand populations harbor very high diversity relative to the invasive populations and are the source of the two main European genetic lineages. One of these two European lineages was in turn the source of at least one of the two main North American genetic clusters of invasive *P. antipodarum*, located in Lake Ontario. The other widespread North American group had a more complex origin that included the other European lineage and two New Zealand clusters. Altogether, our analyses suggest that just a small handful of clonal lineages of *P. antipodarum* were responsible for invasion across continents. Our findings provide critical information for prevention of additional invasions and control of existing invasive populations and are of broader relevance towards understanding the establishment and evolution of asexual populations and the forces driving biological invasion.

### **Keywords**

Approximate Bayesian Computation, SNP,  $Potamopyrgus\ antipodarum$  , invasive route, clonal, genetic diversity

### Introduction

Biological invasions – non-native species that harm native ecosystems and biodiversity (reviewed in Mallez & McCartney, 2018) – are a worldwide problem. Humans have increased the rate of invasions by facilitating the transport of organisms outside of their native range (Nentwig, Bacher, Kumschick, Pyšek, & Vilà, 2018). Genetic variation within the founding and descendent populations is a critical determinant of whether a potential invasion is successful, with high variation more likely to be associated with success (Estoup & Guillemaud, 2010; Estoup et al., 2016). Invasive populations are predicted to lose most of their genetic variation via founder effects during the colonization process (Dlugosch & Parker, 2008), raising the question

of how these invasions succeed (reviewed in Estoup & Guillemaud, 2010; Dlugosch et al., 2015; Estoup et al., 2016). One potential solution to the barrier that low genetic variation should pose to successful invasion could be provided by multiple distinct introductions of colonists (Dlugosch & Parker, 2008; Kolbe et al., 2004). While multiple introductions are indeed associated with invasion success, there are also multiple examples of successful invasions in the face of low genetic diversity (reviewed in Estoup & Guillemaud, 2010; Dlugosch et al., 2015).

These seemingly conflicting results highlight the fact that the relationship between genetic diversity and invasion success is not straightforward (recently comprehensively reviewed in Dlugosch et al., 2015; Estoup et al., 2016), with mechanisms like admixture, genotype-by-environment interactions, or purging of harmful alleles coming into play during the invasion process. A recent demonstration of the complicated relationship between diversity and invasion success is provided by the growing body of evidence suggesting that worldwide invasions might often be catalyzed by successful invasive populations rather than directly via the native range ("bridgehead effect"; Lombaert et al., 2010; e.g., Reed et al., 2020, Wylie et al., 2020). Recent advancements in population genomics resources and tools now allow us to reconstruct invasion routes, enabling us to address key questions regarding the identity of source populations and how genetic variation is influenced by the colonization process (Estoup & Guillemaud, 2010; Dlugosch et al., 2015; Estoup et al. 2016). This information can then be applied in turn to characterize whether and how evolution plays a role in the driving successful invasions (Dlugosch & Parker, 2008; Keller & Taylor, 2008; Estoup et al., 2016).

The wide availability of genomic resources overcomes limitations posed by earlier marker types (e.g., microsatellites) with respect to, for example, reconstructing evolutionary relationships among populations (Mc-Cartney, Mallez, & Gohl, 2019). In parallel, continued development of analytical methods like Approximate Bayesian Computation (ABC) analysis provides inferences that are useful in untangling complex scenarios often seen in invasive history (Estoup & Guillemaud, 2010). This information offers new insights into the specific mechanisms by which species introductions occur, which can be leveraged towards establishing best practices to prevent and control invasions.

Potamopyrgus antipodarum is a fresh- and brakish water prosobranch snail that has become widely known as a model organism for the evolution of sex, characterized by frequent coexistence between obligately sexual and obligately asexual individuals in its native range in New Zealand (Lively, 1987). Known colloquially as the "New Zealand mud snail" (NZMS) in the invasive range, *P. antipodarum* has successfully invaded every continent except Antarctica and Africa (Alonso & Castro-Díez, 2012; Collado & Collado, 2014). These snails were first discovered in England in 1859 (Ponder, 1988). Over the 19th century, NZMS spread rapidly throughout western and central Europe (Hamada et al., 2013). The New Zealand mud snail is now, >160 years post-invasion, considered one of the worst alien species in Europe (Nentwig et al., 2018). The introduction of *P. antipodarum* to Australia is estimated to have occurred at a similar time frame as the European invasion, with the first recorded date of presence in Australia in 1872 (Ponder, 1988). *Potamopyrgus antipodarum* was first introduced to North America in 1987 in the western United States (Taylor, 1987) and colonized the Great Lakes in 1991 (Zaranko et al., 1997). In Japan, *P. antipodarum* was first reported in 1990. These Japanese populations seem to represent an invasion that is distinct from European and US invasions (Hamada et al., 2013). Most recently, *P. antipodarum* was reported in central Chile in 2014 (Collado & Collado, 2014), where it has been extending its range (Collado & Fuentealba, 2020).

Invasive P. antipodarum populations, in contrast to the native range, are exclusively asexual (Alonso & Castro-Díez, 2012). Despite harboring low genetic variation relative to populations in the native range, invasive P. antipodarum have become successful in aquatic ecosystems spanning a wide range of environmental conditions across the globe (Dybdahl & Drown, 2011; Alonso & Castro-Díez, 2012). The high population growth rate that can accompany asexual reproduction is thought to be one of the many reasons that P. antipodarum invasions have been successful (Alonso & Castro-Díez, 2008). Apparent adaptive plasticity of shell morphology and life history traits with respect to local environmental conditions also seems to play an important role in invasion success of P. antipodarum (Kistner & Dybdahl, 2014; Verhaegen, et al. 2018a, in press). Other factors such as ovoviviparity, tolerance for a wide range of chemical and physical conditi-

ons, and release from coevolving parasites might also contribute to the *P. antipodarum* invasion (Alonso & Castro-Diez, 2012; Verhaegen et al., 2018a).

Documented negative effects of these snails on invaded ecosystems are likely driven in large part by their high population growth rates, translating into extraordinarily high density [e.g., 20,000-50,000 individuals/m<sup>2</sup> (Hall et al., 2003, 2006; Alonso & Castro-Díez, 2012)] and with dramatic consequences for invaded ecosystems. Hall et al. (2003) provided a clear example of the potential impact of invasive *P. antipodarum*, demonstrating that the snails consumed 75% of gross primary productivity and dominated nutrient fluxes in a creek in Wyoming, USA. Hall et al. (2006) went on to show that *P. antipodarum* strongly distorted the distribution of secondary production across invertebrate taxa, sequestering much of the available carbon and altering ecosystem function. Alonso & Castro-Díez (2012) argued that *P. antipodarum*'s ability to radically alter ecosystem nutrient fluxes puts the threat it poses on par with such notorious invaders as the zebra mussel.

The genetic background of these invasive P. antipodarum populations has been characterized with mitochondrial DNA (mtDNA) markers, revealing that, as for many other invasive taxa (Estoup et al., 2016), the invasive populations harbor very little diversity relative to the native range (Dybdahl & Drown, 2011; Neiman & Lively, 2004; Verhaegen et al., 2018b). These mtDNA data also indicate that the European invasion is primarily composed of two distinct lineages with haplotypes identical to haplotypes found in New Zealand (Städler et al., 2005; Verhaegen et al., 2018a). Dybdahl and Drown (2011) demonstrated that the picture is broadly similar with respect to the North American invasion and emphasized that this lack of diversity is surprising in light of how rapidly the North American invasive populations have successfully established throughout a wide range of environments. In particular, Dybdahl and Drown (2011) showed that two lineages (US1, US2; defined via a combination of mitochondrial and nuclear markers) predominated in the western and eastern parts of the United States, respectively. The mtDNA haplotypes of lineages US1 and US2 are identical to haplotypes (22 and 37 in Neiman and Lively 2004, respectively) that are also found across Europe (Städler et al. 2005; Verhaegen et al. 2018a) and in New Zealand (Verhaegen et al. 2018b). The other two lineages defined by Dybdahl and Drown (2011), US1a and US3, have genotypes that have to date only been found in the western United States (Dybdahl & Drown, 2011). Dusting (2016) discovered that the dominant western North American lineage US1 and the geographically widespread and relatively common Australian invasive genotype AUS2 share a 22-locus SNP genotype, suggesting a close genetic link between the source populations of these invasions (also see Emblidge Fromme & Dybdahl, 2006). Hershler et al. (2010) used microsatellite genotyping to provide a finer-scale characterization of the genetic diversity of P. antipodarum in the western North American Snake River basin population, finding that this population appeared to have been founded by a few rapidly spreading clones.

Together, these studies demonstrate that *P. antipodarum* is a very successful invader even in the absence of genetic variation. What remains a key question is how the worldwide invasive populations are genetically connected. Reconstruction of the invasion route will illuminate whether invasive P. antipodarum populations represent a tiny and perhaps preadapted subset of native diversity that has experienced strong selection in the invasive range versus an invasion process driven by chance sampling events such as accidental boat transport. This information regarding the sampling process during invasion and the role of chance vs. deterministic processes in invasion is a central component of characterizing the traits and factors that drive invasion success (Keller & Taylor, 2008; Estoup et al. 2016). We used multilocus nuclear SNP markers to characterize the relationships amongst a large set of native New Zealand populations and invasive North American and European populations in the framework of several different population genetics modeling approaches to provide the first population genetic assessment of the relationships of invasive and native P. antipodarum across continents continents [for linguistic simplicity we also refer to New Zealand as continent, and in fact, New Zealand is part of Zealandia, a mostly drowned fragment of Gondwanaland (Mortimer et al. 2017)]. We specifically addressed the key question of whether the successful North American and European P. antipodarum invasions represented similar (or even the same) vs. distinct source populations. Discovering that these populations are similar is significant in suggesting that invasions in North America and Europe are closely linked. This result would also be consistent with a scenario where certain native lineages are more invasive than others and where invasion success does not require high genetic variation. Very distinct

European and American populations will in contrast suggest different source populations for the North American and European invasions, as expected in a case where genetic variation is an important component of invasion success. Regardless of specific outcome, the answer to the question of how closely these continentscale invasions are related to one another will provide important new insights into how the invasion of this destructive snail occurred. A deeper understanding of whether and how successful invasive lineages are connected also helps illuminate the factors that contribute to invasion success, species range expansions, and the persistence of populations that are asexual or harbor low genetic variation.

### Materials and Methods

#### Collection, DNA extraction, and genotyping

Freshwater mudsnail (Potamopyrgus antipodarum) estrogen receptor Identification and expression analysis under exposure to (xeno-)hormones

We included a total of 1617 snails from three continents: 1016 snails (51 sites) from the native range in New Zealand, 445 snails (25 sites) from the invaded range in Europe, and 156 snails from 13 invaded sites in North America (**Table 1**). All our sampling locations were freshwater sites with the exception of the European brackish water site DEBIN (Verhaegen et al., 2018a). The genetic data from the New Zealand snails and most of the European (N = 421, 94.6 %) snails were generated in Verhaegen et al. (2018a,b). Snails from North America and from two additional European populations (Mo and Gb) were newly collected in the summers of 2016 and 2017 and genotyped for this study. These snails were collected alive from submerged rocks and vegetation growing in shallow regions of streams and lakes with kick nets and then housed in spring water and fed freshwater fish food until their transport to the University of Iowa. The snails were then transferred to 10-L tanks filled with carbon-filtered tap water at constant 16°C room temperature and a 12 h day/night cycle. Water was changed 2x/week and dried spirulina and chalk were added as food and calcium sources, respectively.

We sacrificed the snails, used forceps to separate the soft tissue from the shell, and flash froze each snail in liquid nitrogen. We then stored the snails at -80°C until DNA extraction. DNA was extracted from the frozen tissue by first lysing the cells in CHAOS buffer [4 M guanidine thiocyanate, 0.1% N-lauroyl sarcosine, 10 mM Tris hydrochloride (Tris-HCl) at pH 8.0, and 0.1 M 2-Mercaptoethanol in 60 mL of double-distilled water (ddH<sub>2</sub>O); Fukuma et al., 2004] for 24 h. The buffer was emulsified with a volume of phenol-chloroform extraction buffer [100 mM Tris-HCl at pH 8.0, 10 mM ethylenediaminetetraacetic acid (EDTA), and 0.1% sodium dodecyl sulfate in 80 mL of  $ddH_2O$  equal to that of the tissue slurry, an equal volume of phenol chloroform isoamyl alcohol (PCIA, volumetric ratio of 25:24:1), and 5  $\mu$ L of RNase A before vortexing. An equal volume of PCIA was added and centrifuged at 10,000 x g for 30 seconds. Another volume of chloroform was added and then vortexed to emulsify before centrifuging again. We pipetted off the aqueous layer, measured its final volume, and added a tenth of this volume of sodium acetate. We then added to this solution twice this final volume of ice-cold 100% ethanol and stored the samples at -20°C overnight. Next, the samples were centrifuged at 13,000 x g at 4°C for 15 minutes and the supernatant pipetted off. We added 1 mL of ice-cold 70% ethanol before centrifuging at 4°C for 10 more minutes. The supernatant was pipetted off and the samples left to dry overnight before being resuspended in 30 µL Tris low-EDTA buffer (10 mM Tris, 0.1 mM EDTA). The extracted DNA was stored on 96-well plates at -80°C until shipment on dry ice to LGC Genomics (Beverly, UK, www.lgcgroup.com/genomics) for KASP<sup>TM</sup> assay-based genotyping of the same 48 single nucleotide-polymorphism (SNP) markers used in Verhaegen et al. (2018b). Sixteen of these markers were designed by Paczesniak et al. (2013) and 32 by Verhaegen et al. (2018a). The SNP data from the newly genotyped individuals were then merged with the data from Verhaegen et al. (2018ab), for a total N of 1617. Twelve loci were fixed across all individuals and hence discarded. The remaining polymorphic SNPs were the same 36 loci previously found to be variable and in Hardy-Weinberg (H-W) and linkage equilibrium among the New Zealand individuals (Verhaegen et al., 2018b). Forty-five individuals (2.78%)

for which genotyping failed for more than five loci were excluded. Genotypes were then assigned by the infinite alleles model distance index with GenoDive v. 2.0b27 (Meirmans & Van Tienderen, 2004) setting the maximum distance threshold for two genotypes to be considered identical to zero (Rogstad et al., 2002). For all subsequent population genetic analyses, we discarded all but one individual per genotype within each unique population or group in order to avoid treating likely clonemates as independent data points.

#### Genetic distances between sites

We calculated pairwise Slatkin's linearized fixation indices based on  $F_{\rm ST}$  (Slatkin, 1995) as implemented in Arlequin v.3.5.2.2 (Excoffier & Lischer, 2010) to evaluate the genetic relationships among sampling sites by building a neighbour-joining (NJ) tree in PAST v.3.25 (Hammer et al., 2001). Two individuals of the closely related *P. estuarinus*, for which the same SNP loci were previously genotyped by Verhaegen et al. (2018b), served as outgroup to root the NJ tree. We used GenGIS v.2.5.3 (Parks et al., 2013) to geographically map the NJ tree.

#### Clustering

We used clustering methods and discriminant analyses of principal components (DAPC) (Jombart et al., 2010) with the R package *adegenet* v.2.1.1 (Jombart, 2008) to evaluate genetic structure amongst native and invasive *P. antipodarum* populations. We first estimated the number of genetic clusters with a K-means clustering analysis based on the lowest Bayesian information criterion (BIC) (Schwarz, 1978). We then used a DAPC to visualize the relationships among these clusters and to assign the individuals to the clusters *a posteriori*. DAPC first transforms the data using a principal component analysis (PCA) and then identifies clusters through a discriminant analysis (DA). Unlike a STRUCTURE-like approach (Pritchard et al., 2000), a DAPC does not rely on population genetic assumptions and therefore can accommodate populations containing clonal individuals. The geographical distribution of these clusters was then mapped with QGIS v.2.18.24 using the vector maps publicly available from Natural Earth (https://www.naturalearthdata.com/). In order to quantify differences between clusters, we calculated the uncorrected number of mutational changes between pairs of genotypes from DAPC clusters including European and North American individuals with PAUP v.4.0b10 (Swofford, 1998).

#### ABC analyses

Approximate Bayesian Computation (ABC) analyses implemented in DIYABC v.2.1.0 (Cornuet et al., 2014) were used to retrace the invasion route of *P. antipodarum* to Europe and North America. While DIYABC assumes random mating within populations, we followed the lead of previous successful applications of DIYABC to other clonal (e.g., Goss et al., 2014; Zhang et al., 2014) or mixed sexual/asexual systems (e.g., Barres et al., 2012; Janoušek et al., 2016; Taerum et al., 2017). For the native range, we divided the New Zealand individuals into two groups, one per island (North vs. South Island, labelled NZN and NZS). For the invaded range, we took into account our clustering results and previously established mtDNA haplotypes (Verhaegen et al., 2018a) to divide the European snails into two groups (EU14 vs. EU15). Based on the clustering results, the North American samples were divided into two groups (US1 vs. OnBrF). The ABC groups are summarised in Table 2. Three of our 36 polymorphic SNP markers (SNP IDs: comp148591\_c0\_seq1, comp149304\_c0\_seq1, and comp160266\_c0\_seq4) were discarded from all our ABC analyses because they could not be genotyped for all individuals within at least one group. In order to optimize the computational effort, our ABC analyses were performed in a stepwise fashion as proposed by Lombaert et al. (2014) following the chronological order of the introductions into Europe and the US. In a first step, we clarified the most likely invasion scenario of *P. antipodarum* to Europe. In a second step, we tested all plausible invasion scenarios to the US. For this first step, we tested, in one ABC analysis (ABC1), 12 plausible invasion scenarios of P. antipodarum to Europe (Supplemental Fig. 1). This included six scenarios without unsampled "ghost" populations (i.e., unsampled populations from anywhere geographically between the sampled New Zealand

sites and the invaded regions), four scenarios with one ghost population originating from either NZN or **NZS**, and two scenarios with two ghost populations, each originating from a different New Zealand island. As P. antipodarum diversity is highest on the North Island (Neiman & Lively, 2004), and considering that P. antipodarum's five most closely related congeners are confined to the North Island (Haase, 2008), we set **NZN** as the most ancestral group in all scenarios. The unit of time of the historical scenarios in DIYABC is set by the generation time of the study organism (Cornuet et al., 2014). Assuming that the generation time of P. antipodarum is 6-12 months (Larkin et al., 2016; Verhaegen et al., in press; Winterbourn, 1970) and that the introduction of *P. antipodarum* to Europe happened about 180 years ago (Ponder, 1988; Smith, 1889), we set the divergence time of our European groups to a range of 180 to 360 generations. This interval takes into account the uncertainty of the date of the actual introduction and the documented variation in generation time. Posterior probabilities for the scenarios were estimated with a logistic regression on the 1%of simulated datasets that were closest to the observed dataset (Cornuet et al., 2008, 2010). We allowed the software to perform a linear discriminant analysis on the summary statistics before the regression in order to reduce the computation time (Cornuet et al., 2014; Estoup et al., 2012). The most likely scenario was then selected based on these probabilities ensuring that the 95% confidence intervals of competing scenarios did not overlap (Cornuet et al., 2008).

In a second step, we used the most likely scenario of the first step as a base to identify the most likely invasion scenarios of *P. antipodarum* to North America. It was not computationally feasible to analyse all possible scenarios against each other at once, so we divided the 36 scenarios into four different ABC analyses. In a first analysis (ABC2), we tested 16 invasion scenarios without the presence of ghost populations (Supplemental Fig. 2). In a second analysis (ABC3), nine scenarios including a ghost population originating from NZN and two scenarios with two ghost populations originating from NZN and NZS, respectively, were tested against each other (Supplemental Fig. 3). In ABC4, we tested nine scenarios including a ghost population originating from NZS (Supplemental Fig. 4). In ABC5, we then tested the most likely scenario of each of the first three analyses (ABC2-4) against one other. In the last analysis, ABC6, based on the clustering results, we tested the most likely scenario of ABC5 against two scenarios with either US1 originating both from EU15 and NZN, or from EU15 and NZS. As P. antipodarum is estimated to have been introduced to North America about more than 30 years before our sampling (Bowler, 1991; Taylor, 1987; Zaranko et al., 1997), we set the divergence time parameter for North American populations to a range from 35 to 70 generations, again considering the uncertainty of the actual arrival of P. antipodarum as well as the variation in generation time. As the introduction date of P. antipodarum to Australia is estimated to be around the same time as to Europe (Ponder, 1988), we set the divergence time for the ghost populations to be equal or younger than the divergence time of the European populations.

All of our ABC analyses except for ABC2 (671,093 simulations/scenario, over six times the minimum required simulation number) and ABC3 (944,373 simulations/scenario, over nine times the minimum required number of simulations) were performed with one million simulated datasets per scenario, the optimum number recommended for DIYABC. We used fewer than one million simulations for ABC2 and ABC3 because of computational constraints. The default minimum allele frequency criterion for SNP data was used, namely the Hudson's algorithm (Hudson, 2002). We calculated the following summary statistics: 1) per population, the proportion of loci with null gene diversity (i.e., the proportion of monomorphic loci), mean and variance of the gene diversity across polymorphic loci (Nei, 1987), and the mean gene diversity across all loci; 2) per population pair, the proportion of loci with null  $F_{\rm ST}$  distance (Weir & Cockerham, 1984), the mean and variance across loci of non-null  $F_{\rm ST}$  distances, and the mean across loci of  $F_{\rm ST}$  distances. These summary statistics are used to discriminate among scenarios and to compute posterior distributions of parameters. A bottleneck event was included after all divergence events. The effective population size of all tested populations was set to be less than or equal to the size of NZN and the effective size of all bottleneck populations was set to be less than or equal to the populations that emerged from these bottlenecks. The divergence time of NZS from NZN was set to be greater than all the other divergence times present in our scenarios. The goodness of fit of the final chosen scenario was tested through a PCA comparing the prior and posterior distributions of the summary statistics to the observed data, as implemented in DIYABC

("Perform model checking").

### Results

#### Population genetic analyses and NJ tree

We detected a total of 338 SNP genotypes in our dataset; 309 of these genotypes were collected from New Zealand, 20 from Europe, and nine from North America. At 16 of the 89 sites all individuals shared the same genotype (three New Zealand sites, five European sites, and eight North American sites). For sites harbouring more than one genotype, the genetic diversity was low to moderate, with values for Ho varying between 0.016 and 0.486 ( $0.279 \pm 0.038 - \text{mean} \pm \text{SD}$ ), and for Hs between 0.016 and 0.353 ( $0.213 \pm 0.026$ ) (Table 1). The NJ tree based on Slatkin's linearized fixation indices between pairs of sites (Supplemental Table 1) is given in Fig. 1 and geographically mapped in Fig. 2. European and North American clades were separated and scattered across the entire tree. In only one case - the sample from Ontario- was the closest relative of one of these invasive genotypes not a genotype or clade from New Zealand. This Ontario sample emerged from a European clade.

#### Clustering

In order to retrace the invasion route of P. antipodarum, we first assessed the genetic structure by means of DAPC clustering. With an overall a posteriori assignment probability of 93.9% (individual a posteriori probabilities are found in **Supplemental Table 2**), individuals were grouped into 15 different clusters (**Fig. 3**). All 15 clusters included individuals from New Zealand (four clusters exclusively from the North Island, seven clusters from the South Island, and the remaining four clusters from both islands), 11 of these clusters contained only New Zealand individuals. Clusters 1 and 2 consisted of New Zealand individuals and five North American individuals from Sn and one from Ml, respectively. Individuals from all US sites except On, European sites DEBIN, Mo, NL1, NL2, and two New Zealand individuals from NZ52 and one from NZ64 were found in cluster 15. Individuals from the North American site On, one individual from US site BrF, another individual from NZ64, and all European sites except DEBIN formed cluster 14. The European individuals from cluster 15 were pooled together as group **EU15** for the ABC analyses and the European snails from cluster 14 as **EU14 (Table 2**). The geographical distribution of these clusters is given in **Fig. 4** and bar plots with the number of individuals per continent within each cluster in **Supplemental Fig. 6**.

The variation within and between the four clusters including individuals from the invaded range was given by the uncorrected number of mutational changes between pairs of their genotypes. Overall, genotypes from these clusters differed from each other by 1 - 36 mutations (**Supplemental Table 3**). Genotypes within clusters 1, 2, 14, and 15 differed from each other by 1 - 20, 1 - 9, 1 - 23, and 1 - 14 mutations, respectively. All but one individual from the American site On and one individual from BrF shared genotype 324 with one another and were pooled together as the ABC group **OnBrF** (**Table 2**). Genotype 324 was also shared with individuals from **EU14** present at sites Gb and Mo. One remaining individual from site On harbored a genotype that differed only by one mutation from genotype 324 and was therefore included in **OnBrF**. There was relatively high genotypic variation present among American individuals of **US1** (**Table 1**), which harboured seven genotypes from cluster 15 differing from each other by one to four mutations, one genotype (337, at least six mutations from New Zealand genotypes) from cluster 1, and one genotype (332, at least five mutations from New Zealand individuals) from cluster 2. The most common and geographically widespread **US1** genotype (325) was also found in one individual from **EU15** (site Mo), and differed by at least eight mutations from European or New Zealand genotypes. The other **US1** genotypes from cluster 15 were at least six mutations away from the two other **EU15** genotypes.

#### **ABC** analyses

All five initial ABC analyses (ABC1-5) had a most likely scenario clearly distinct from competing scenarios with a posterior probability > 97%. The most likely invasion route to Europe (ABC1, scenario 6, Supplemental Fig. 1) had EU14 originating from NZN and EU15 from NZS and did not include ghost populations [0.9773 (0.9724,0.9821) – posterior probability (95% confidence interval)]. For the three preliminary ABC analyses (Supplemental Fig. 2 to 4) testing the invasion route to North America, scenarios 6 [ABC2, 0.9985 (0.9975, 0.9994)], 11 [ABC3, 0.9996 (0.9994, 0.9998)], and 9 [ABC4, 0.9996 (0.9995, 0.9998)] emerged as the most likely scenarios for their respective analysis. These three most likely scenarios were tested against each other in ABC5 as scenarios 1, 2, and 3, respectively (Supplemental Fig. 5). Here, scenario 1 was the most likely invasion scenario [0.9996 (0.9995, 0.9998)], in which **OnBrF** originated from EU14, US1 from EU15, and the assumption of ghost populations was not necessary. The probability of scenario 1 was nevertheless low  $[0.1625 \ (0.1536, 0.1715)]$  when tested against the two admixture scenarios in the final ABC6, with the most likely scenario as US1 originating from EU15 and NZN [scenario 2, 0.6193 (0.6069,0.6317)], and the second most likely scenario as US1 originating from EU15 and NZS [scenario 3, 0.2182 (0.2080,0.2283)] (Fig. 5). The final invasion scenario retained by our ABC analyses was summarized and mapped in **Fig. 6**. Posterior probabilities and confidence intervals for all tested scenarios are given in **Supplemental Table 4**. Despite the unambiguity of the posterior probabilities for the scenario choice, the final scenario 2 of ABC6 showed a poor goodness of fit (Fig. 7). The posterior distribution of the summary statistics formed a distinct cloud that did not contain the observed data in the PCA plot checking the model. This discrepancy may be due to the limitations that ABC methods face with respect to asexually reproducing organisms.

### Discussion

We used genotyping of nuclear SNPs from a large and geographically diverse sample of *P. antipodarum* from invaded sites in North America and Europe and the native range in New Zealand to characterize its invasion route across continents. In particular, the combination of traditional and Bayesian molecular analyses allowed us to resolve outstanding key questions regarding potential direct connections between geographically distinct invasions. Together, these data complement and extend earlier genetic and genomic work in *P. antipodarum* (Städler et al., 2005; Dybdahl & Drown, 2011; Verhaegen et al., 2018a) by demonstrating conclusively that major components of the worldwide NZMS invasion were driven by just a few asexual lineages. We also showed for the first time that the European invasion was seeded by introductions from several different New Zealand populations and that this European invasion in turn is a major source of the invasion of North America. Together, these data provide a qualitative advance forward in suggesting that a bridgehead scenario (Lombaert et al., 2010) is an apt framework for this destructive global invasion. More broadly, these results are of relevance to understanding other invasions of asexual organisms and/or invasions with relatively low genetic variation, and provides critical information on genetic diversity and invasion sources, central in any invasion control strategy.

#### From where did the European snails originate?

European snails were divided in two distinct genetic groups **EU14** and **EU15**, based on the clustering results of our DAPC (clusters 14 and 15, respectively). Individuals of **EU14** and the Great Lakes shared the mtDNA haplotype 22 (Dybdahl & Drown, 2011; Neiman & Lively, 2004; Verhaegen et al., 2018a). This haplotype was, however, absent from our New Zealand samples (Verhaegen et al., 2018b), but reported in foregoing studies from both islands [e.g., in Lakes Waikaremoana, Gunn, and Te Anau (Neiman & Lively, 2004; Neiman et al., 2011)]. Using SNP markers, the ABC enabled us to retrace the origin of **EU14** to the North Island. While the only New Zealand-sourced individual from cluster 14 was collected on the South Island (site NZ64), our NJ tree showed that clades from the North Island harboured the closest relatives to the European sites of cluster 14, indicating the high dispersive potential of *P. antipodarum*. Genotyping New

Zealand individuals harbouring haplotype 22 from both New Zealand islands will provide an important next step, allowing us to test if the North Island was indeed the origin of **EU14**. As all ABC scenarios testing the origin of **EU14** from unsampled ghost populations were rejected and because no Australian *P. antipodarum* of haplotype 22 has been found (Dusting, 2016), we can state with some confidence that individuals of **EU14** were directly introduced from New Zealand.

By contrast, the ABC analysis indicated that **EU15** originated from New Zealand's South Island. For this analysis, New Zealand individuals of mtDNA haplotype 37 from both the North and South Island were present within our tested dataset (Verhaegen et al., 2018b). A few individuals from DAPC cluster 2 were also found in New Zealand, but only on the South Island (sites NZ52 and NZ64). Invasive scenarios including ghost populations as the origin of **EU15** were rejected by the ABC analysis. Because haplotype 37 is also found in Australia (Dusting, 2016), a useful validation step would come from the inclusion of Australian individuals in future analyses. However, because the introduction of *P. antipodarum* to Australia was reported several decades after the European introduction (Brazier, 1871; Hubendick, 1950; Ponder, 1988), we do not expect Australia to emerge as a stepping stone between New Zealand and Europe in subsequent studies.

#### What is the source(s) of the North American invasion?

First, the DAPC analyses revealed that the North American snails were represented predominantly by the same two genetic clusters present in Europe (clusters 14 and 15). This result suggests that P. antipodarum of at least two different geographic and/or genetically distinct sources successfully colonized North America. Lake Ontario (ABC group **OnBrF**, all from cluster 14) P. antipodarum were genetically distinct from the other North American sites (ABC group **US1**, predominantly from cluster 15). Individuals from two other eastern US sites (Mr and PA) were also part of this widespread North American cluster 15. One individual from group **OnBrF** was found in the Bear River in Idaho (site BrF), more than 2500 km west of the other P. antipodarum with the same genotype in Lake Ontario. This result again indicates the high potential of *P. antipodarum* for passive dispersal across large distances (Hubendick, 1950). Additional genotyping of more individuals from the BrF site will be needed to determine whether this western North America snail with an apparently "eastern" genotype is common in this region, providing important information about the connectivity across North American freshwater sites. The general pattern that seems to emerge - that individuals from the Great Lakes and from western US sites constitute two very distinct, different clonal lineages - aligns with the results of earlier analyses of mitochondrial DNA variation (Dybdahl & Drown, 2011; Neiman, et al., 2010, 2011). In a qualitative departure from these earlier studies, we also found that this "western" US haplotype is in fact not limited to western North American sites. Indeed, snails from two eastern North American sites (Mr and PA) shared the same genetic cluster with the majority of the genotypes from the western sites.

We were able to use our data to provide a conclusive answer to the long-standing question of the origin of the *P. antipodarum* invasion of North America. All our analyses also revealed that the Great Lakes group of North American *P. antipodarum* represented by cluster 14 originated in Europe. This result is reinforced by the fact that all individuals from Lake Ontario shared a SNP genotype with individuals from European sites Gb and Mo and differed by one to eight mutations from the other **EU14** genotypes. Individuals of **EU14** (except from the newly sampled and unsequenced sites Mo and Gb) were already known to share the same mtDNA haplotype 22 with individuals from the Great Lakes (Dybdahl & Drown, 2011; Verhaegen et al., 2018a), but so did some New Zealand populations (Neiman & Lively, 2004; Neiman et al., 2011). The use of nuclear SNP markers allowed us to exclude the latter as the source of the North American Great Lakes *P. antipodarum*.

The origin of the North American *P. antipodarum*, represented by ABC group **US1**, is more complex. The ABC suggested that **US1** originated both from Europe [European DAPC cluster 15 (ABC group **EU15**)] and from New Zealand's North Island (ABC group **NZN**). These results are consistent with the DAPC clustering. The NJ tree also showed that New Zealand clades from both islands as well as two European clades (DEBIN and DERUG) are closely related to or ancestral to the **US1** clade. The distribution of **EU15** is much more

restricted than **EU14** and included our only brackish-water site, DEBIN. Individuals from **EU15** (except unsequenced site Mo) shared mtDNA haplotype 37 (Verhaegen et al., 2018a), the same haplotype also found in the western USA (Dybdahl & Drown, 2011). The most common genotype of **US1**, genotype 325, was also shared with one individual from the **EU15** site Mo. However, within this same cluster 15, the other genotypes from **EU15** differed by at least eight mutational steps from those of **US1**. It is difficult to reconcile so many changes accumulating at third-codon positions in protein-coding nuclear DNA sequences with an introduction of asexual snails to western North America only 30 years ago (Bowler, 1991). Accordingly, if **US1** *P. antipodarum* indeed originated from **EU15**, it seems to us more likely that their introduction to North America occurred substantially earlier than reported or that these western North America snails originated from European or New Zealand sites not included in our survey.

The second origin of North American **US1** snails would have been directly from New Zealand, where individuals of haplotype 37 are also present (Neiman & Lively, 2004; Neiman et al., 2010, 2011), including individuals from our dataset (Verhaegen et al., 2018b). A few individuals from **US1** were also part of clusters absent in Europe but present in New Zealand. Twenty-four percent of the individuals from Idaho's Snake River (site Sn) were included in cluster 1, a cluster otherwise restricted to the North Island of New Zealand. One individual from Idaho's Malad River (Ml) was part of cluster 2, present on both New Zealand islands. **US1** snails in Idaho harbored relatively high genetic variation (see also BrF discussed above). Because the first reported North American *P. antipodarum* were found in the Snake River (Bowler, 1991), Idaho, it is possible that this variation could have arisen since invasion. If this variation did not arise *in situ*, this assemblage must be due to additional dispersal, possibly through transportation by birds (see Zielske et al., 2017). This latter scenario, however, is in contradiction to the ABC analyses rejecting ghost populations.

To summarise, our North American sites were divided into two distinct genetic groups: one in Lake Ontario, and another spread across the continent. All our analyses pointed towards Europe as the invasion source of *P. antipodarum* from the Great Lakes: some identical SNP genotypes were even shared between some European snails and those of Lake Ontario. The origin of the other widespread North American group is more complex: our analyses pointed towards both Europe and New Zealand as likely sources.

#### ABC analyses in the context of asexual reproduction

Recent developments in genomic technologies and sophisticated analytical methods allow us to indirectly retrace historical invasion routes of species that would have otherwise remained invisible (Estoup & Guillemaud, 2010). ABC methods provide a qualitative step forward by enabling the testing of invasion scenarios including unsampled "ghost" populations or specific stochastic events (e.g., admixture events). Like most population genetic analysis frameworks, ABC analyses specifically designed for asexually reproducing organisms are yet to be developed. We addressed this limitation of the ABC approach by excluding all but one member of each clone within each ABC group (e.g., Janoušek et al., 2016; Taerum et al., 2017). Even so, the relatively poor goodness of fit of the final invasion scenario that was selected by this analysis suggests that even this modification might fall short in this instance by simulating the genetic genealogies using algorithms that assume recombination (e.g., in our case the Hudson's algorithm, suitable for SNP data, Hudson, 2002) (Cornuet et. al, 2014). Nevertheless, ABC is still often used to test evolutionary scenarios involving systems that are asexual (e.g., wheat aphids, Zhang et al., 2014), mixed sexual/asexual (e.g., plant pathogens, Goss et al., 2014) or that alternate between asexual and sexual reproduction (e.g., trees, Lander et al., 2011; or plant fungi, Barrès et al., 2012; Janoušek et al., 2016; Taerum et al., 2017). While some of these studies also removed all but one member of a clone (e.g., Janoušek et al., 2016; Taerum et al., 2017), even this correction might not be necessary: Barrès et al. (2012) compared the ABC analyses using clone-corrected vs. uncorrected datasets and found no differences in outcome.

*Potamopyrgus antipodarum* is very similar to many of the invasive taxa focused on in these other studies in that invasive lineages are asexual while native populations harbor both asexually and sexually reproducing individuals, but is distinct in the high genetic diversity of clonal *P. antipodarum* in the native range (e.g., Paczesniak et al., 2013) relative to invasive populations. By contrast, the genetic diversity of invasive clonal

populations in these previous studies that did not seem to experience poor model-fitting was high relative to our invasive *P. antipodarum* populations (e.g., Goss et al., 2014; Zhang et al., 2014), with the former possibly mimicking conditions expected in diverse and outcrossing populations. This possibility suggests that a useful next step might be a simulation-based study investigating the robustness and appropriateness of DIYABC under different levels of clonal diversity.

Despite the poor fit of our final ABC model, the ABC approach is still of use in terms of providing distinct and complementary information relative to non-Bayesian analyses. For example, comparisons of the outcomes of these different approaches revealed similar outcomes of the NJ tree and ABC analyses with respect to the invasion route of **OnBrF**, but different outcomes for **US1**. In this latter case, while the NJ tree suggested New Zealand as origin of **US1**, the ABC instead pointed to both Europe and New Zealand as origin. How do we reconcile the different outcomes of these two analyses? We can begin by considering how the analyses are structured: in particular, the NJ tree is generated from pairwise genetic distances between sampling sites, while the ABC uses genotype data. Sampling sites can contain individuals from different genotype clusters, which can result in apparently contradictory inferences when using methods that assess relationships among sampling sites (i.e., NJ tree) vs. individuals (i.e., ABC). Although our data structure likely violates the ABC modeling assumptions, we cannot yet estimate to which extent the results, which appear intuitively reasonable (Fig. 4), are actually misleading. For example, group US1 contained individuals from clusters 1 and 2 (dominating in New Zealand) and Cluster 15, which probably arrived via Europe and the eastern United States. This pattern suggests that US1 has a more complex origin involving several sources. This in fact turned out to be the most likely scenario. With this in mind, and given that violations of ABC assumptions are not necessarily detrimental (Barres et al. 2012; Goss et al., 2014; Zhang et al., 2014), we should not blindly discard these inferences until simulations have tested the robustness of DIYABC analyses.

#### Conclusion and perspectives

Here, we used characterization of the genetic connections across invasive populations of a destructive snail and the relationships of these populations to native counterparts to determine the source of the invaders and how the invasion process has affected population genetic diversity. The answers to these questions are a critical component of understanding invasion success (Keller & Taylor, 2008; Dlugosch et al., 2015; Estoup et al., 2016). More broadly, these data also address important unresolved issues involving the evolutionary potential of asexual lineages and populations with low genetic variation.

Our analyses revealed that the native range of New Zealand is the source of the two main European genetic lineages of *P. antipodarum*. One of these European lineages was the unique source of the North American group of the Great Lakes, whereas the other European lineage was the source, in combination with two New Zealand clusters, of the second main widespread Nothern American cluster. In other words, besides demonstrating low genetic variability on a fine genetic scale across two continents, we found that only a few clonal lineages of *P. antipodarum* are responsible for the contential North American and European invasion.

There are multiple and often non-mutually exclusive potential explanations for invasion success in the context of low genetic diversity (reviewed in Estoup et al., 2016). First, a compelling "null" hypothesis is the absence of an adaptive challenge in the native range, which can occur, for example, in situations where the environments in the native and invaded range are similar. The niche modeling work of Loo et al. (2007) does seem to exclude this possibility, demonstrating that invasive P. antipodarum thrive in environments found outside New Zealand. Another hypothesis that must be considered is that relatively low variation in molecular markers does not always translate into relatively low variation in ecologically relevant traits (e.g., conversion of non-additive variation during a bottleneck; reviewed in Neiman & Linksvayer, 2005). Recent studies do reveal substantial genetic variation for fitness-relevant traits in invasive P. antipodarum (Neiman & Krist, 2016; Levri et al., 2017), indicating that this scenario deserves more attention. It is also possible that these successful invasive lineages of P. antipodarum can compensate for low genetic variation through phenotypic plasticity (reviewed in Estoup et al. 2016; e.g., Verhaegen et al., 2018a). Taken together, our new data and these earlier studies hint that relevant and heritable variation - even in the almost total absence of molecular marker-based estimates of genetic diversity - was likely present in the New Zealand and, subsequently, bridgehead European populations that seeded future invasions.

This hypothesis can be evaluated by including further unsampled potential native source populations more recently invaded regions like Japan and Chile (Hamada et al., 2013; Collado, 2014; Collado & Fuentealba, 2020) as well as comparing the phenotypic means and variances of native vs. invasive clones (e.g.,Keller & Taylor, 2008; Neiman & Krist, 2016; Levri et al., 2017). These studies will provide a powerful test of the role - if any - of genetic and phenotypic variation in driving initial vs. later invasions. Further study of *P. antipodarum* from these perspectives will also help illuminate how asexual lineages and populations with low genetic diversity succeed in new environments.

### Acknowledgments

We thank the Iowa Science Foundation, the University of Iowa Center for Global & Regional Environmental Research, and the Dewey Stuit Fund for Undergraduate Research for the collection of *P. antipodarum* at the North American sites BrB, BrF, Co, Gr, LC, Md, Ml, Pn, and Sn. We thank Dr. Amy Krist and Dr. Ed Levri for supplying samples from North American sites Pc and Mr, On, and PA, respectively. Several anonymous reviewers provided very useful and constructive feedback. This study was funded by the Research Training Group 2010 RESPONSE grant of the Deutsche Forschungsgemeinschaft (DFG) for the European and New Zealand data, and by the Iowa Science Foundation for the North American data.

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### **Data Accessibility Statement**

The genetic data used in this study is given in the Supplemental material, as Supplemental Files 1 to 3.

### Author contributions

- Manuscript: GV, CD, MN, MH, JDW
- Design, concept: GV, MN, MH
- Fieldwork of new samples: JDW, MN
- DNA extraction of new samples: JDW
- Genetic data of Europe and New Zealand: GV, MH
- Analyses: GV, CD, MH

## Tables

**Table 1** Sampling sites and results of the genetic diversity summary statistics as implemented in Genodive analyses. Standard genetic results (Ho and Hs) were not calculated for the asexual populations in the invaded range, neither for New Zealand populations that had all their individuals sharing the same SNP genotype.

Footnote:  $^{+}ABC = Approximate Bayesian Computation, <math>^{++}Ho = observed heterozygosity, ^{SS}Hs = expected heterozygosity.$ 

Acror	n <b>Glo</b> un- try	Location	Date (month year)	Lati- tude	Longi- tude	Ν	ABC+ group	Ho+ <b>H</b> s§	Number of unique genotypes
NZ01	New Zealand	Whatawhata, West Waikato	Febru- ary 2016	S 37° 47' 7 09"	E 175° 4' 7 26"	20	NZN	0.3 0.2	7
NZ03	New Zealand	Whatawhata	Febru- ary 2016	S 37° 47' 2 3"	E 175° 4' 10 4"	20	NZN	0.34 0.23	8
NZ04	New Zealand	Bridal Veil Falls	Febru- ary 2016	S 37° 54' 31 17"	E 174° 53' 45.4"	20	NZN	0.23 0.22	8
NZ07	New Zealand	McLaren Falls	Febru- ary 2016	S 37° 48' 52 57"	E 176° 2' 48.02"	20	NZN	0.2 0.18	7
NZ08	New Zealand	McLaren Falls	Febru- ary 2016	S 37° 49' 3 02"	E 176° 2' 49.19"	20	NZN	0.32 0.19	2
NZ10	New Zealand	Kaniwhaniwha reserve	Febru- ary 2016	5.02 S 37° 55' 12.70	49.19 E 175° 4' 52.88"	20	NZN	0.38 0.28	6
NZ14	New Zealand	Wairere falls	Febru- ary	12.79 S 37° 44'	52.88 E 175° 51'	20	NZN		1
NZ19	New C Zealand	rown track, Karangahake	Febru- ary	13.89 S 37° 25'	52.5 E 175° 43'	20	NZN	0.08 0.08	19
NZ21	New N Zealand	orth of Kaimai-Mamaku Forest Park	Febru- ary 2016	40.17 S 37° 28' 20.41"	$E 175^{\circ} 46'$	20	NZN	0.28 0.21	4
NZ23	New N Zealand	forth of Kaimai-Mamaku Forest Park	Febru- ary 2016	20.41 S 37° 27' 51 84"	44.07 E 175° 46'	20	NZN	0.09 0.09	18
NZ24	New N Zealand	orth of Kaimai-Mamaku Forest Park	Febru- ary 2016	S 37° 27' 47.00"		20	NZN	0.32 0.25	2
NZ27	New Zealand	Waitomo	Febru- ary 2016	41.03 S 38° 15' 57 2"	±3.07 E 175° 4' 37.47"	20	NZN	0.27 0.24	7
NZ29	New Te Zealand	e waihou walkway to blue spring	Febru- ary 2016	S 38° 1' 59 8"	E 175° 49' 31.82"	20	NZN	0.17 0.17	11
NZ31	New Zealand	Wainui falls track, Tata beach	Febru- ary 2016	S 40° 50' 36 16"	E 172° 56' 9.52"	20	NZS	0.31 0.28	7
NZ32	New M Zealand	IcShane road, Tata beach	Febru- ary 2016	S 40° 49' 43 24"	E 172° 56' 44 49"	20	NZS	0.38 0.28	5
NZ37	New Zealand	Collingwood, McDonald Footbridge	Febru- ary 2016	S 40° 40' 52 41"	$E 172^{\circ}$ 40' 26.01	20	NZS	0.38 0.22	2
NZ39	New Zealand	Takaka	2010 March 2016	S 41° 4' 49 3"	$E 172^{\circ}$ 44'	20	NZS	0.29 0.24	2
NZ41	New Zealand	Salmon farm, Takaka	March 2016	$18^{\pm 5.5}_{S 40^{\circ}}_{53'}_{0 73''}$	$E 172^{\circ} 46' 1.51''$	20	NZS	0.36 0.26	7
NZ42	New Zealand	Dry road, Rakopi	March 2016	S 40° 36' 33.84"	E 172° 34' 10.05"	20	NZS	0.39 0.25	5

Acronym	Individuals the group contains
NZN	Individuals from New Zealand North Island
NZS	Individuals from New Zealand South Island
EU14	European individuals included in DAPC cluster 14
EU15	European individuals included in DAPC cluster 15
OnBrF	American individuals included in DAPC cluster 14, i.e. all individuals from site On and one from BrF
$\mathbf{US1}$	American individuals included in DAPC clusters 1, 2 and 15, i.e. individuals from all North American sites ex

Table 2 Groups of individuals used in the Approximate Bayesian Computation analyses

### Figure legends

**Figure 1** Neighbour-joining tree based on pairwise Slatkin's linearized  $F_{ST}$  values and rooted by *Potamopy-rgus estuarinus* (= pink). Blue = New Zealand sites; orange = North American sites; green = European sites. Corresponding information for the acronyms can be found in Table 1.

**Figure 2** Geographical mapping of the neighbour-joining tree. Blue = New Zealand sites; orange = American sites; green = European sites; pink, *Potamopyrgus estuarinus* (outgroup).

Figure 3 Scatterplot of the first two axes of the discriminant analysis of principal components (DAPC) showing the relationship of 15 clusters. Ellipses represent inertia ellipses. Clusters in dark blue contain exclusively New Zealand individuals; light blue = cluster 1; pink = cluster 2; green = cluster 14; orange = cluster 15. The DAPC was built by maintaining the first 36 PCs and six discriminant functions.

Figure 4 Geographical distribution of the 15 DAPC clusters. Clusters in dark blue only contained individuals from New Zealand and are therefore not distinguished from each other on this map.

**Figure 5** The three competing invasion scenarios of the final Approximate Bayesian Computation (ABC6) analysis with their posterior probabilities and 95% confidence intervals in square brackets. The selected scenario, scenario 2, is highlighted in bold. Bottleneck events represented by dotted lines. The timeline is not to scale. For the constitution of the different ABC groups, see Table 2.

**Figure 6** Simplified map-based representation of our current picture of the *P. antipodarum* global invasion. Non-black line colours represent cluster membership for EU14, EU15, OnBrF, and US1. Lines are drawn to roughly represent known shipping routes most likely to have been the source of *P. antipodarum* colonization. The dotted black lines connecting New Zealand to Tasmania and Tasmania to South Australia represent likely invasion routes, though not tested directly here. We have also included the recent Chilean and Japanese invasions on the map, though the genetic data needed to infer invasion routes are not yet available.

**Figure 7** Principle component analysis (PCA) for checking the goodness of fit of the most likely scenario 2 of the final Approximate Bayesian Computation analysis ABC5. The PCA compares the prior and posterior distribution of the summary statistics, and the observed data set. The variance explained by each principle component (PC1 and PC2) are given in brackets.

### Supplemental figures legends

**Supplemental Fig. 1** Invasion scenarios to Europe tested in the Approximate Bayesian Computation analysis ABC1. Bottleneck events are indicated with a lower case "b", ghost populations as "NA".

**Supplemental Fig. 2** Invasion scenarios to the USA without presence of ghost populations tested in the Approximate Bayesian Computation analysis ABC2. Bottleneck events are indicated with "b".

**Supplemental Fig. 3** Invasion scenarios to the USA including a ghost population originating from the population NZN (New Zealand North Island), and scenarios with two ghost populations originating from NZN and NZS (New Zealand South Island), respectively, tested in the Approximate Bayesian Computation analysis ABC3. Bottleneck events are indicated with "b"; ghost populations as "NA". The scenarios are given in the historical model parameterization code format of DIYABC v.2.1.0 (Cornuet et al., 2014).

**Supplemental Fig. 4** Invasion scenarios to the USA including a ghost population originating from the population **NZS** (New Zealand South Island), tested in the Approximate Bayesian Computation analysis ABC4. Bottleneck events are indicated with "b"; ghost populations as "NA". The scenarios are given in the historical model parameterization code format of DIYABC v.2.1.0 (Cornuet et al., 2014).

Supplemental Fig. 5 The three competing invasion scenarios of the Approximate Bayesian Computation analysis ABC5 with their posterior probabilities and 95% confidence intervals in square brackets. The selected scenario, scenario 1, is highlighted in bold. Unsampled "ghost" populations are indicated in grey; bottleneck events represented by dotted lines. The timeline is not to scale. For the constitution of the different ABC groups, see Table 2.

**Supplemental Fig. 6** Distribution of the individuals from New Zealand (blue), Europe (green), and North America (orange) within the 15 clusters. Only one individual per unique SNP genotype was kept per sampling site for the clustering analysis.

### Supplemental tables legends

Supplemental Table 1 Pairwise Slatkins linearised  $F_{\rm ST}$  values between sampling sites.

**Supplemental Table 2** A priori and a posteriori cluster assignment for all the individuals, as well as a posteriori cluster assignment probability.

**Supplemental Table 3** Uncorrected number of mutational changes between pairs of genotypes from DAPC clusters including European and American individuals (excel sheet 1); number of individuals for each genotype per continent and DAPC cluster (excel sheet 2)

**Supplemental Table 4** Posterior probabilities and 95% confidence intervals for all tested scenarios in the Approximate Bayesian Computation analyses ABC1-6.

### Supplemental files legends

Freshwater mudsnail (Potamopyrgus antipodarum) estrogen receptor Identification and expression analysis under exposure to (xeno-)hormones

**Supplemental File 1** Raw SNP results for all 1617 collected snails. Excel sheet 1 and 2 = results for the European snails genotyped in Verhaegen et al. (2018a), sheet 3 = results for the New Zealand snails genotyped in Verhaegen et al. (2018b), sheet 4 = results for the US and European snails genotyped in this study, sheet 4 = summary of polymorphic and fixed SNP markers among continents. Corresponding information for the population acronyms can be found in Table 1. More information on the SNP IDs can be found in the Supplementary Table 2 of Verhaegen et al. (2018b).

**Supplemental File 2** GENEPOP file used for the genetic population analyses, using the same 36 polymorphic SNP loci as in Verhaegen et al. (2018b). All but one individual per genotype within each population were discarded in order to avoid treating likely clonemates as independent data points. Adenine bases were coded as "01", cytosine as "02", guanine as "03", thymine as "04', and unknown bases as "00".

Supplemental File 3 Input file with our SNP data for the ABC analyses ABC2 – 6 conducted with DIYABC v.2.1.0 (Cornuet et al., 2014). Three of our 36 polymorphic SNP markers (SNP IDs: comp148591\_c0\_seq1,

comp149304\_c0\_seq1, and comp160266\_c0\_seq4) were discarded from all our ABC analyses because they could not be genotyped for all individuals within at least one group. All but one individual per genotype within each ABC group were discarded in order to avoid treating likely clonemates as independent data points. For ABC1, only the New Zealand and European groups were used.





Figure 2: Geographical mapping of the neighbour-joining tree. Blue = New Zealand sites; orange = American sites; green = European sites; pink, *Potamopyrgus estuarinus* (outgroup).



Figure 3: Scatterplot of the first two axes of the discriminant analysis of principal components (DAPC) showing the relationship of 15 clusters. Ellipses represent inertia ellipses. Clusters in dark blue contain exclusively New Zealand individuals; light blue = cluster 1; pink = cluster 2; green = cluster 14; orange = cluster 15. The DAPC was built by maintaining the first 36 PCs and six discriminant functions.



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