

Invasion genetics of vendace (*Coregonus albula* (L.)) in the Inari-Pasvik watercourse: revealing the origin and expansion pattern of a rapid colonization event

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Introduction

Introductions and invasions of exotic species represent a problem of global extent (Vitousek et al. 1996; Williamson 1996; Olden and Rooney 2006). Severe ecological effects may arise from invasions or possible interactions between climate change and invasions (Sandlund et al. 1999; Mooney and Hobbs 2000; Driscoll et al. 2012), and the consequences may also be associated with huge economical costs (Moyle 1986; Mack et al. 2000; Pimentel et al. 2000). During recent decades, numerous studies

Abstract

Species invasions can have wide-ranging biological and socio-economic effects and are generally unwanted by legislation. Identification of the source population as well as the ecology and genetics of both the invader population and the receiving community is of crucial importance. The rapid invasion of a small coregonid fish vendace (*Coregonus albula*) in a major northern European subarctic watercourse has resulted in a labile ecological situation in the receiving community. The ecological impact of the invasion has been thoroughly documented, but the genetics of the invasion remains to be explored. We analyzed the genetic diversity and divergence patterns among the two possible source populations from southern Finnish Lapland and three colonist populations within the Inari-Pasvik watercourse using ten microsatellite loci in order to (i) identify the most likely source of the invasion, (ii) reveal the dispersal pattern and genetic structure of the secondary expansion, and (iii) to investigate whether the initial introduction and the secondary expansion were associated with founder effects. We revealed that repeated translocation of vendace from Lake Sinettjärvi into a tributary lake of L. Inari in 1964–1966 is the most plausible source for the invasion. Both the initial introduction and the secondary expansion were found not to be associated with significant founder effects. The secondary expansion followed a stepping stone pattern and the source and colonist populations of this expansion have undergone rapid genetic divergence within a period of 15–35 years (ca. 8–17 generations). The rapid divergence may be contributed to lack of gene flow among the source and colonist populations due to the extensive hydroelectric damming in the watercourse. Multiple introductions and substantial genetic variation in combination with the boom-and-bust population development of the species thus likely counteracted the founder effects as well as fueled the rapid establishment and expansion of this species within the Inari-Pasvik watercourse.

have addressed the ecological effects of biological invasions, particularly with respect to the consequences for the receiving communities. Negative impacts include imperilment of native species (Allan and Flecker 1993; Peterson et al. 2004; Bøhn et al. 2008), alterations of community structure and natural biodiversity (Williamson 1996; Parker et al. 1999), and a worldwide biotic homogenization (Rahel 2000, 2002; Olden and Poff 2004; Olden and Rooney 2006). However, species introductions may also be seen as large-scale ecological experiments, and provide unique insight into ecological interactions as

well as population biology and genetics (Sakai et al. 2001; Shea and Chesson 2002; Sax et al. 2007; Davis 2009).

The genetic consequences and outcome of an invasion invent for the invaders and the colonist populations, that is, the correlation between propagule pressure, genetic variability of the invaders and the resulting genetic variability in the colonist populations, are not trivial. Colonist populations, that is, the populations descending from introduction events, may rapidly develop differences in morphology and phenotypes (Facon et al. 2008; Ward et al. 2012) and in ecology (Sandlund 1992; Sakai et al. 2001; Amundsen et al. 2012) as compared to the source population, but the genetic component is often unknown (e.g., Blanchet 2012). Colonist populations can give rise to secondary expansions, that may cover large geographical scales (Facon et al. 2008; Brown and Stepien 2009; Reusch et al. 2010). Such demographic expansions propel divergent adaptive pressures due to exploration of new environments (Ghalambor et al. 2007), which again may promote rapid changes in life history of the invader (Dlugosch and Parker 2008a; Amundsen et al. 2012; Gutowsky and Fox 2012). While most studies have found lower genetic variation in the colonist compared to the source populations (Henry et al. 2009; Ayres et al. 2010; but see Dlugosch and Parker 2008b for review), others show no loss of genetic variation (Novak and Mack 2005; Stepien et al. 2005; Wares et al. 2005). Some studies even find increased genetic variation in the colonist populations when they originate from multiple introductions promoting genomic admixture (Golubtsov et al. 1993; Kolbe et al. 2004; Lavergne and Molofsky 2007). The fraction of genetic variation transferred from the source populations to the invaders depends on propagule pressure (Lockwood et al. 2005; Colautti et al. 2006; Simberloff 2009), dispersal mode (Hewitt 1996; Ibrahim et al. 1996; Wilson et al. 2009), and/or route of invasion (Estoup and Guillemaud 2010). The altered selection pressure applied when individuals explore new geographical ranges also represents challenges for the invader (Lee 2002; Suarez and Tsutsui 2008). Thus, it is the combined effect of these factors that determine whether the invaders are successful in their colonization and proliferation.

The present study concerns the introduction and subsequent invasion of an exotic fish species (vendace, *Coregonus albula*, Fig. 1) into the subarctic Inari-Pasvik watershed (Norway, Finland, and Russia). The first introduction occurred in 1956, when some vendace fry were translocated for stocking purposes from Lake Kelujärvi in central Finland to the Inari Hatchery, from where they subsequently escaped into Lake Inari (Mutenia and Salonen 1992; see also Fig. 2). The second introduction occurred in 1964–1966 when vendace fry were translocated from Lake Sinettäjärvi to a small lake in the catchment area of Lake



Figure 1. Invading vendace (*Coregonus albula*) from the Pasvik-Inari watercourse, Norway (photo: K. Ø. Gjelland).

Inari. The first vendace were observed in Lake Inari in 1973. By the early 1980s, a vendace population had established throughout the lake (Mutenia and Salonen 1992), increasing to a peak abundance in 1989 (Salonen 1998, 2004). During this period a downstream invasion of vendace apparently occurred to lakes in the Pasvik watercourse, the outlet river from Lake Inari, where the species was observed for the first time in 1989 (Amundsen et al. 1999). The vendace were observed in L. Vaggatem for the first time in 1991 and in L. Skrukkebukta in 1993, and within few years the invader became an important pelagic fish species in lakes in the Pasvik watercourse (Bøhn et al. 2004, 2008). Severe ecological consequences have been documented for the receiving system, including changes in biodiversity and species composition, food web dynamics and ecosystem functioning (e.g., Bøhn and Amundsen 1998, 2001; Amundsen et al. 2003). The vendace have during the invasion entered a typical fluctuating ‘boom-and-bust’ development (Salonen et al. 2007), resulting in a destabilized aquatic ecosystem.

Although documentation of the ecological impact of the vendace invasion is accumulating, the genetics of the invasion remains to be addressed. A recent study revealed reproductive isolation between vendace in L. Inari and a downstream lake (Amundsen et al. 2012), which initiated this more detailed genetic study of the system. Because we hold detailed temporal and ecological information about the invasion process (see e.g., Mutenia and Salonen 1992; Gjelland et al. 2007; Bøhn et al. 2008) and have the special opportunity to precisely identify the source of the invasion, the system offers an excellent model system to assess ongoing genetic and evolutionary changes. We also expect the downstream invasion to be highly influenced by the seven hydropower dams that exist in the watercourse (Amundsen et al. 2012). The dams only allow unidirec-

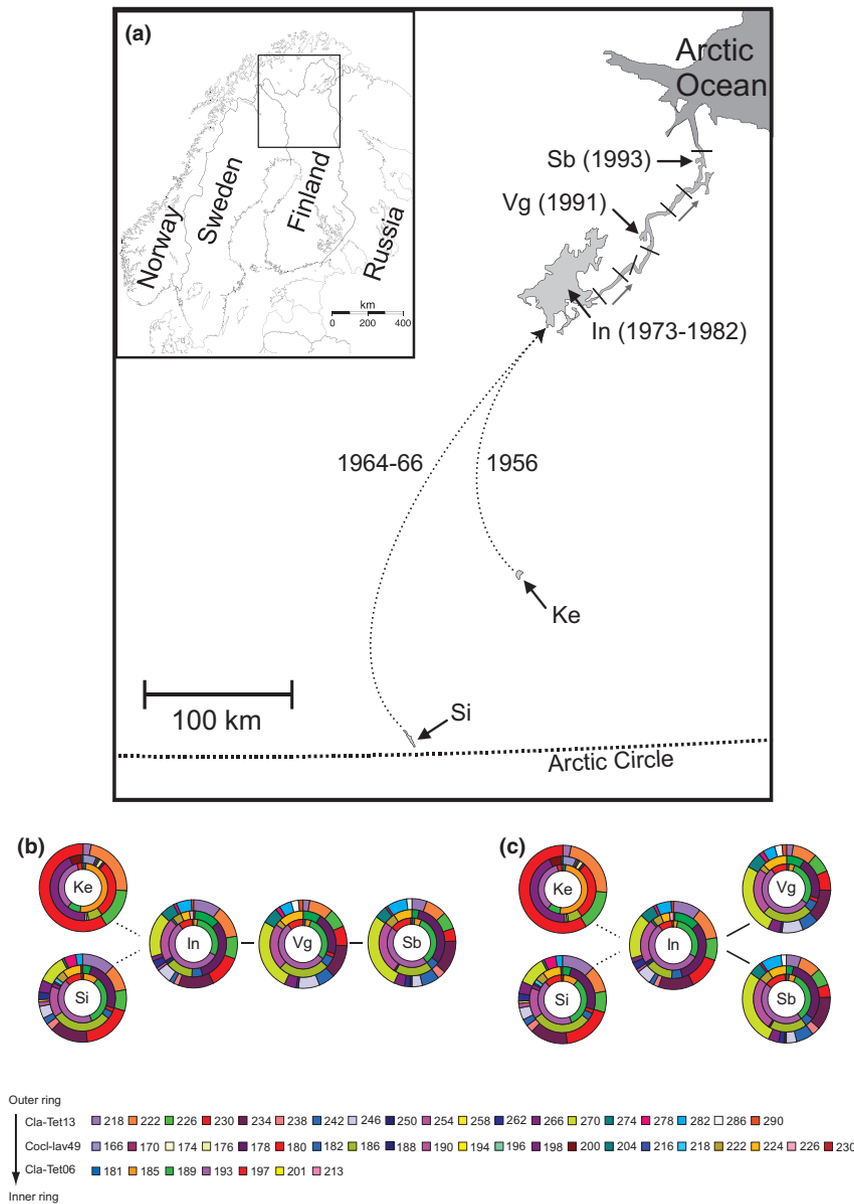


Figure 2. Origin and sample locations (a) and molecular genetic variation (b, c) of vendace in northern Fennoscandia. In (a) light gray arrows show the direction of drainage. Also, note that hydroelectric dams (indicated by black lines in the map) are located between the introduced L. Inari population (In) and the populations within the secondary expansion (Vg and Sb) limiting downstream gene flow and ruling out any upstream gene flow. The schematic illustration of the invasion process following a stepping stone (b) and normal dispersal pattern (c) are illustrated with the corresponding pie charts of the relative allelic frequencies of three representative microsatellite loci (ClaTet13, Cocl-lav49, and ClaTet06). Dashed lines indicate man-made translocations (L. Sinettäjärvi and L. Kelujärvi to L. Inari) and full black lines delineate the natural secondary expansion for the invasion. Below are the color codes for the alleles per locus. Sample locations are abbreviated according to Table 1.

tional downstream gene flow that would be limited or absent during normal river flow conditions, as the fish larvae or adults would need to pass through dam turbines (see also Amundsen *et al.* 1999, 2012). Hence, potential factors that may limit the successful colonization in the system may relate both to founder effects and to unidirectional gene flow. Here, we used ten microsatellites to explore the genetics of the introduction and subsequent downstream migration of vendace in the watercourse by comparing the two potential source populations, the introduced population, and two downstream colonist populations originating from the secondary expansion. The main objectives of the study were to (i) identify the most likely source of the invasion, (ii) reveal the dispersal pattern and

genetic structure of the secondary expansion, and (iii) to investigate whether the initial introduction and the secondary expansion were associated with founder effects.

Materials and Methods

Study lakes and sample collection

To identify the most likely origin of the invasion, vendace were sampled from the introduced population in L. Inari, northern Finland, and the two potential source populations, L. Sinettäjärvi and L. Kelujärvi, central Finland (Table 1; Fig. 2). Vendace were furthermore sampled from the colonist populations in L. Vaggatem and

Table 1. Locations and codes samples included in the genetic analysis of the northern European vendace invasion (Fig. 2), with latitude and longitude (position), sample date, water drainage, lake size and height of location in water course, date of translocation (for L. Kelujärvi and L. Sinettäjärvi to L. Inari) or first observation on locality (for L. Inari, L. Vaggatem and L. Skukkebukta), secchi depth, and sample size (*N*).

Lake	Code	Position	Sample date	Water course (Ocean)*	Area (km ²)	Height (m)	Year (Observed)	Secchi depth (Max depth, m)	<i>N</i>
L. Kelujärvi	Ke	67°28'N, 27°4'E	14.11.2008	Kemijoki (BS)	9	187	1956	2 (10)	52
L. Sinettäjärvi	Si	66°35'N, 25°25'E	25.10.2008	Kemijoki (BS)	9	97	1964–1966	3 (40)	58
L. Inari	In	69°10'N, 27°55'E	30.10.2008	Inari-Pasvik (AO)	1102	118	1973	5 (95)	55
L. Vaggatem	Vg	69°13'N, 29°14'E	09.09.2008	Inari-Pasvik (AO)	34	52	1991	3 (30)	57
L. Skukkebukta	Sb	69°33'N, 30°7'E	16.09.2008	Inari-Pasvik (AO)	7	21	1993	6 (38)	54

*BS, Baltic Sea; AO, Arctic Ocean.

L. Skukkebukta in the Norwegian part of the Inari-Pasvik watercourse to investigate the genetics of the secondary expansion. All samples were collected in the autumn 2008 near or at the spawning grounds and consisted of >92% ripe individuals. There are seven hydropower dam constructions located below L. Inari (Fig. 2). More detailed descriptions of L. Inari and the Pasvik watercourse are given by Mutenia and Salonen (1992) and Amundsen et al. (1999), respectively.

Microsatellite DNA amplification and genotyping

Genomic DNA was extracted from gill filaments by E-Z96 Tissue DNA Kit (OMEGA Bio-tek, Norcross, GA) following the manufacturer instructions. The individuals were genotyped at 10 microsatellite loci (Table S1) arranged in three multiplex polymerase chain reactions (PCR) following a previously described protocol (Præbel et al. in press). The PCR products were separated on an ABI 3130 XL Automated Genetic Analyzer (Applied Biosystems, Foster City, CA) and alleles scored in the GeneMapper 3.7 software (Applied Biosystems). After the first validation of the genotypes, 3–4% of the individuals within each population were re-extracted and rerun at all ten loci. The genotypes resulting from the initial run and the rerun were manually compared for all individuals to rule out miss-scoring of alleles. If any doubt occurred in this comparison the samples were re-extracted and rerun at all loci to obtain a consensus genotype. The samples were finally screened for abnormalities (null alleles, scoring errors, etc.) in the software MICRO-CHECKER 2.2.3 (Van Oosterhout et al. 2004), using 1000 bootstrap replications to generate the expected homozygote and heterozygote allele size difference frequencies.

Statistics

The within population genetic variation indices; number of alleles (N_A), expected (H_e) and observed (H_o) heterozygos-

ity, and the fixation index (F_{IS}) were estimated in GenAEx 6.41 (Peakall and Smouse 2006; Table S1). Deviations from Hardy–Weinberg equilibrium (HWE) for each population and locus and linkage disequilibrium (LD) among loci and among loci over all populations were tested by exact tests (Guo and Thompson 1992) using GENEPOP 4.0 (Rousset 2007). The tables of pair-wise *P*-values from the LD and HWE tests were corrected for multiple comparisons by sequential Bonferroni corrections (BFCs) following Rice (1989). Allelic (N_{RA}) and private allelic richness (N_{RPA}) per population were determined, accounting for differences in sample-sizes, using the rarefaction procedure for the smallest sample size (100 genes) as implemented in the software HP-RARE 1.0 (Kalinowski 2005).

To infer which of the two initial vendace stockings (L. Sinettäjärvi or L. Kelujärvi) in L. Inari that led to successful colonization of the Inari-Pasvik watercourse, a neighbor-joining tree was build using Nei et al. (1983) genetic distance (D_a) and nodes were tested for robustness by 1000 bootstraps using Populations 1.2.32 (Langel-la 2005) and viewed in TREEVIEW (Page 1996). A sample of native vendace ($N = 20$) obtained from northern Germany was used to assist the clustering of the neighbor-joining tree. Genetic divergence between the possible source populations and the introduced L. Inari population as well as among the colonist populations were estimated by pair-wise F_{ST} (Weir and Cockerham 1984) values and tested for statistical significance (10,000 permutations) using ARLEQUIN 3.5.1.2 (Excoffier and Lischer 2010). The table of *P*-values for the pair-wise F_{ST} values was corrected for multiple comparisons by BFCs following Rice (1989). Bayesian clustering as implemented in STRUCTURE 2.3.2 (Pritchard et al. 2000; Hubisz et al. 2009), was used to provide another estimate of origin and population structure of the data. We used a model assuming admixture and correlated allele frequencies between *K* populations (Burn-ins of 100,000 replications and 300,000 Markov chain Monte Carlo (MCMC) replicates). Sampling locations were used as a priori information to assist the structuring (the LOCPRIOR model)

as recommended for weak signals of structuring (Hubisz *et al.* 2009). All runs were replicated 10 times at each $K = 1-5$ to confirm consistency of log-likelihood probabilities. The most likely (highest $\ln \Pr(X|K)$) grouping was visualized using STRUCTURE HARVESTER (Earl and vonHoldt 2012). Structuring of the vendace populations were also estimated by principal component analysis (PCA). The ordination of the individuals was performed in the program GenAlEx 6.41 (Peakall and Smouse 2006). Finally, we investigated whether shared private alleles could be identified in pair-wise comparisons of the source L. Kelujärvi and/or L. Sinettäjärvi and the introduced population (L. Inari), as well as among populations within the watercourse which would support any link found in the population structuring approach. The rarefaction procedure implemented in ADZE 1.0 (Szpiech *et al.* 2008) were used to estimate the pair-wise shared private allelic richness across loci for the possible source populations and the introduced population (L. Inari) and for the secondary expansion within the water course (i.e., between L. Inari-L. Vaggatem, L. Inari-L. Skrukkebukta, and L. Vaggatem-L. Skrukkebukta). All estimates were performed using a standardized sample size corresponding to the smallest sample (100 genes).

We tested whether the initial stocking and the subsequent invasion was associated with founder effects by testing for genetic bottlenecks among the populations using the software BOTTLENECK 1.2.02 (Cornuet and Luikart 1996). Population bottlenecks will cause a temporarily imbalance in the mutation-drift equilibrium, where additions of new alleles via mutation are balanced by the loss of alleles via drift (Luikart and Cornuet 1998). We utilized this assumption to identify situations where the actual sample heterozygosity exceeds a permuted equilibrium heterozygosity as expected under a mutation-drift equilibrium. If the actual heterozygosity exceeds the equilibrium heterozygosity it is indicative of a recent population bottleneck (Cornuet and Luikart 1996). We used 1000 coalescent simulations and assumed a two-phased model of mutation (TPM) and the more conservative step-wise mutation model (SMM). The statistical significance of the deviations at equilibrium and observed heterozygosities were tested with Wilcoxon signed-rank tests. We also tested for genetic signatures of founder effects following the invasion and the secondary expansion, by estimates of changes in the effective population size (N_e). Changes in the effective population size will indicate the relative contribution of genetic drift (Hedrick 2000). N_e was estimated for all populations using OneSamp 1.1 (Tallmon *et al.* 2008). This software uses approximate Bayesian computation to estimate variance N_e from summary statistics that are related to N_e . We used prior

upper and lower bounds for N_e of 2–1000 and 10,000 replications to generate the 95% credible intervals. Finally, we correlated the pair-wise F_{ST} 's and the difference in expected heterozygosity between the source population and Inari-Pasvik populations (ΔH_e) with geographic distance and years from translocation to occurrence. This was performed to reveal spatial and temporal patterns of the mechanism underlying the founder event and to estimate the time of the possible expansion from L. Inari downstream the Pasvik watercourse. We expected increased differentiation (pair-wise F_{ST} 's) and difference in genetic diversity (ΔH_e) the longer the distance and time since divergence. The correlation of genotype data and geographical distance (physical divergence) from the source population to each of the populations in the Inari-Pasvik watercourse were compared with the corresponding correlations for the three possible expansion times. The time from introduction to the founding of the L. Inari population was in all estimates set to 8 years, whereas the expansion time from L. Inari downstream the watercourse was tested for initial sighting of vendace larvae (1973), total cover of lake (1980), and peak (1989) (Table S2). We used partial Mantel tests as implemented in GenAlEx 6.41 (Peakall and Smouse 2006), to test these scenarios, using 9999 permutations to obtain significance.

Results

Genotyping, validation, and quality control of genotypic data

We did not identify any mismatch between the original individual multilocus genotypes and the re-extracted 3–4% replicates within the present dataset. Heterozygote deficits were indicated by MICRO-CHECKER at BWF1 (L. Vaggatem, L. Skrukkebukta), C2-157 (L. Skrukkebukta), Cocl-lav06 (L. Sinettäjärvi, L. Inari, L. Vaggatem, and L. Skrukkebukta) Cocl-lav10 (L. Sinettäjärvi, L. Inari, L. Vaggatem, and L. Skrukkebukta), Cocl-lav49 (L. Skrukkebukta), ClaTet06 (L. Kelujärvi), ClaTet13 (L. Sinettäjärvi and L. Skrukkebukta) all indicated as caused by the presence of null alleles. To test whether the loci with potential null alleles may affect the results the STRUCTURE analyses were performed as described in the Material and Methods but without the loci showing the most heterozygote deficits across populations (BWF1, Cocl-lav06, and Cocl-lav10). Neither the number of inferred clusters nor the population structure was found to differ when comparing the results with and without these loci (Figs. S1 vs. 4). Given the relatively few loci used in the study we therefore maintained all loci in the full analysis to ensure statistical power.

Genetic variation

The standard indices of within population genetic variation are given in Tables 2 and S1. We discerned 124 alleles among the ten microsatellite loci assayed in the five studied vendace populations, with a within population variation of 47 to 89 alleles. Mean number of alleles (N_A) per locus per population varied from 2 to 25 and expected (H_e) and observed heterozygosity (H_o) per locus per population varied from 0.019 to 0.916 and 0.019 to 0.895, respectively. All populations, except L. Kelujärvi, showed significant departures from HWE associated with heterozygote deficits. Linkage disequilibrium was identified for 4 out of 45 pair-wise locus tests across populations but all returned non-significant after sequential BFC. The individual locus tests displayed 18 out of 50 and 7 out of 50 significant deviations from HWE before and after BFCs, respectively (Table S1), and 2 to 9 significant LD's within each population (only three significant after BFC, all in L. Sinettjärvi).

The allelic richness were almost double in L. Sinettjärvi, L. Inari, L. Vaggatem, and L. Skrukkebukta ($N_{RA} = 7.9$ – 8.8) compared to L. Kelujärvi ($N_{RA} = 4.6$) (Table 2). Private allelic richness varied between $N_{RPA} = 0.11$ – 0.97 with the lowest N_{RPA} found in the lower Pasvik river lakes. H_e and H_o were also higher in L. Sinettjärvi, L. Inari, L. Vaggatem, and L. Skrukkebukta ($H_e = 0.596$ – 0.614 ; $H_o = 0.506$ – 0.546) compared to L. Kelujärvi ($H_e = 0.400$; $H_o = 0.369$) (Table 2).

Identifying the source for the vendace invasion and population structure within the secondary expansion

The pair-wise F_{ST} estimates of genetic differentiation between the two possible source populations (L. Kelujärvi/L. Sinettjärvi) and L. Inari suggest that L. Sinettjärvi was the source population as this population pair display the lowest, although significant, differentiation (L. Kelujärvi vs. L. Inari, $F_{ST} = 0.158$, $P < 0.0001$; L. Sinettjärvi vs. L. Inari, $F_{ST} = 0.011$, $P = 0.0027$; Table S3). L. Kelujärvi moreover appeared to be the most

isolated population of the five populations studied ($F_{ST} = 0.141$ – 0.158 ; $P < 0.0001$). Within the Inari-Pasvik watercourse, the introduced population (L. Inari) was significantly different from the populations from the secondary expansion (L. Vaggatem and L. Skrukkebukta), whereas the populations from L. Vaggatem and L. Skrukkebukta could not be significantly discriminated (Table S3). The sequential BFCs did not change the significance level for any of P -values from the pair-wise F_{ST} estimates. The PCA plot revealed groupings of (i) L. Kelujärvi and (ii) L. Sinettjärvi, L. Inari, L. Vaggatem, and L. Skrukkebukta (Fig. 3). Each axis, PC1 and PC2, explained 26.6% and 24.4% of the total variation, respectively.

The Bayesian clustering revealed that the individuals could be partitioned into two genetic clusters ($K = 2$; $\ln Pr(X|K) \pm SD = -7209 \pm 1$; Fig. S2), separating L. Kelujärvi into one and L. Sinettjärvi, L. Inari, L. Vaggatem, and L. Skrukkebukta into the other cluster (Fig. 4). The combined approach of a neighbor-joining tree with Nei et al. (1983) genetic distance (D_a) provided support for L. Sinettjärvi being the most likely source for the vendace in the Inari-Pasvik watercourse. Moreover, the neighbor-joining tree also provided support for divergence between the founder population of L. Sinettjärvi and the introduced population in L. Inari (bootstrap support of 100%) as well as between L. Inari and the populations from the secondary expansion (L. Vaggatem and L. Skrukkebukta, 98%), which is also supported by the low, but significant F_{ST} values. Finally, the Bayesian clustering and the neighbor-joining tree grouped L. Vaggatem and L. Skrukkebukta together supporting the non-significant F_{ST} value estimated between these populations.

A higher mean number of shared private alleles were found within the L. Sinettjärvi-L. Inari pair compared to the L. Kelujärvi-L. Inari pair (Fig. 5a). The mean number of private alleles shared by the L. Kelujärvi-L. Sinettjärvi pair resembled that of the L. Kelujärvi-L. Inari pair, supporting the conclusion that L. Sinettjärvi is the source population for the invasion. For the secondary expansion the pair-wise comparison of mean number of shared pri-

Table 2. Summary genetic statistics of the vendace populations included in the study.

Sample	F_{IS}	H_e	H_o	N_A	N_{RA}	N_{RPA}	TPM	SMM	N_e
Ke	0.0772	0.400	0.369	47	4.6	0.79	0.084	0.003 (d, 0.002)	55 (38–153)
Si	0.1133*	0.614	0.544	89	8.6	0.97	0.322	0.019 (d, 0.009)	234 (130–988)
In	0.0870*	0.597	0.546	89	8.8	0.73	0.193	0.007 (d, 0.003)	250 (150–746)
Vg	0.0849*	0.596	0.546	81	7.9	0.11	0.625	0.019 (d, 0.009)	150 (93–422)
Sb	0.1664*	0.607	0.506	79	8.4	0.58	0.275	0.003 (d, 0.002)	200 (112–703)

The coefficient of inbreeding (F_{IS}), genetic diversity estimates (H_e/H_o), total number of alleles (N_A), mean allelic (N_{RA}), and private allelic richness (N_{RPA}). The P -values for the Bottleneck test are given (TPM and SMM) and significant values (SMM) were only found significant for heterozygote deficit (denoted with d and the corresponding P -value for the exact test). The effective population sizes (N_e) with credible intervals are given. Sample codes as in Table 1.

*Significant heterozygote deficit (positive F_{IS}) estimated by Hardy–Weinberg exact tests.

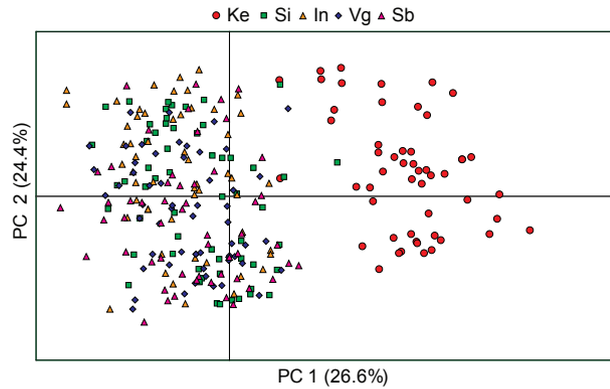


Figure 3. Principal component analysis (PCA) plot of the genetic structuring among the five vendace populations. PC1 and PC2 explain 26.6% and 24.4% of the total variation, respectively. Sample abbreviations as in Table 1.

vate alleles show that the L. Inari-L. Vaggatem pair share fewer private alleles than the L. Vaggatem-L. Skrukkebukta pair, with the L. Inari-L. Skrukkebukta pair sharing the lowest number of private alleles (Fig. 5b). This suggests a stepping-stone dispersal where L. Inari vendace colonized L. Vaggatem and the population in L. Vaggatem subsequently founded the population in L. Skrukkebukta (Fig. 2b). Some specific alleles (e.g., allele 270 at ClaTet13, allele 190 at Cocl-lav49, but see Fig. 2b) were also observed only in the L. Sinettäjärvi, L. Inari, L. Vaggatem, and L. Skrukkebukta populations and display an expanding pattern, while others (e.g., allele 189 at ClaTet06) were present in all populations but in much higher/lower frequencies in L. Kelujärvi compared to the other four lakes (Fig. 2b).

Founder effects and patterns of spatial and temporal genetic variation

None of the populations showed significantly higher expected heterozygosity (H_e) than equilibrium heterozygosity (H_{eq}) in the analysis of bottlenecks assuming a TPM. Similar results were revealed when assuming a SMM. All estimates of effective population sizes were associated with overlapping CIs (Table 2). However, the estimate of L. Sinettäjärvi and L. Inari did not differ in magnitude, supporting that the introduction of vendace into the system was associated with limited founder effects (drift). A decreasing trend in effective population size (although all CIs overlapped) was revealed from L. Inari to L. Skrukkebukta suggesting an increased influence of drift and restricted gene flow during the expansion.

All spatial and temporal correlations of genotypes were found to be significant (Table 3). The period of peak abundance of vendace (1989) in L. Inari also correlate the best with patterns of genetic expansion (ΔH_e), suggesting

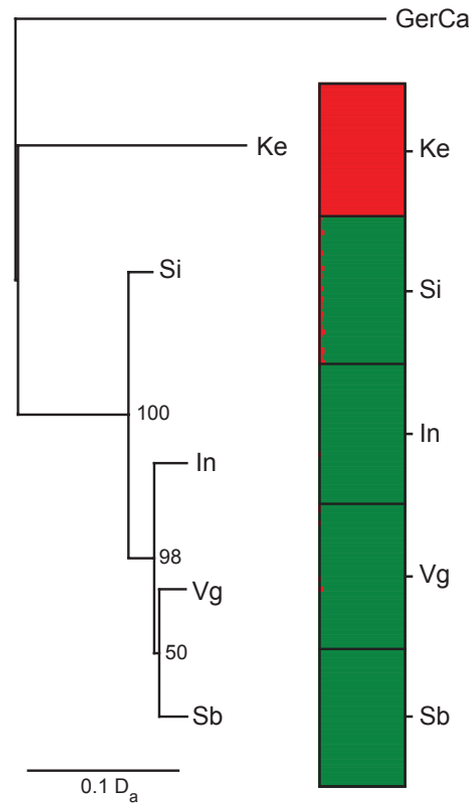


Figure 4. Genetic structure among L. Kelujärvi (Ke), L. Sinettäjärvi (Si), L. Inari (In), L. Vaggatem (Vg), and L. Skrukkebukta (Sb) as revealed by a neighbor-joining trees using Nei et al. (1983) genetic distance (D_a) and Bayesian clustering of all individuals using STRUCTURE (Hubisz et al. 2009) assuming two genetic clusters of individuals ($K = 2$). Only bootstrap resampling percentages above 50 are shown for the neighbor-joining tree. In the STRUCTURE analysis black lines separate individuals from different sampling sites (labeled right) and each individual is represented by a thin horizontal line, which is partitioned into K -colored segments representing individual's estimated membership fractions in K clusters. The absolute mean values of $\ln \Pr(X|K)$ are plotted for $K = 1-5$ in supplementary Fig. S2.

that the secondary expansion downstream the watercourse occurred late in the invasion process and that it was associated with genetic constrains. This is also in accordance with the observation of vendace in catches in Pasvik; vendace were first observed in the upper Pasvik in 1989, and then successively downstream the watercourse (Vaggatem in 1991 and Skrukkebukta in 1993; Amundsen et al. 1999).

Discussion

Primary source of the Inari-Pasvik vendace invasion

There were two possible sources for the introduction and subsequent downstream invasion of vendace in the Inari-

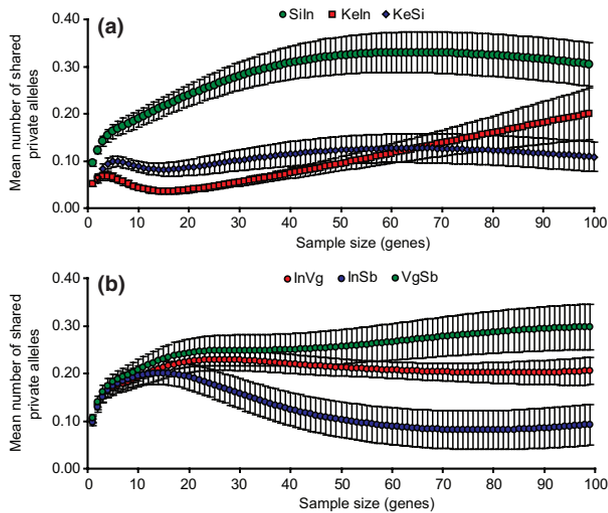


Figure 5. The mean number of shared private alleles private to location pairs of L. Kelujärvi-L. Inari (KelIn), L. Sinettjärvi-L. Inari (SilIn), and L. Kelujärvi-L. Sinettjärvi (KeSi) (a) and within the Inari-Pasvik watercourse (L. Inari-L. Vaggatem, InVg; L. Inari-L. Skrukkebukta, InSb; and L. Vaggatem-L. Skrukkebukta, VgSb) (b) as a function of standardized sample sizes (in genes). Error bars represent \pm SEM across the 10 microsatellite loci.

Table 3. Spatial and temporal correlations of pair-wise differences (F_{ST}) and differences in expected heterozygosity (ΔH_e) using partial Mantel tests.

	F_{ST}		ΔH_e	
	R_{xy}	P	R_{xy}	P
Geographical distance	0.548	0.041	-0.987	0.040
Time _{Initial_observation}	0.778	0.045	-0.623	0.040
Time _{Total_coverage}	0.682	0.045	-0.817	0.042
Time _{Peak_abundance}	0.483	0.042	-0.912	0.041

The coefficient of regression (R_{xy}) and the corresponding P -value (9999 permutations) are given. Geographical distances and expansion times used for correlations are given in Table S2. ΔH_e , Pair-wise difference in heterozygosity between L. Sinettjärvi and the populations within the Inari-Pasvik watercourse.

Pasvik watercourse (Mutenia and Salonen 1992; Salonen 1998). Multilocus microsatellite data and several statistical approaches clearly identified L. Sinettjärvi vendace as the most likely founder population of Inari-Pasvik vendace. The low, but significant, pair-wise F_{ST} value found between the vendace populations of L. Sinettjärvi and L. Inari indicates that these populations have diverged recently. In comparison, the F_{ST} value between L. Kelujärvi and L. Inari was high and represents the typical level of genetic divergence reported between other

native postglacial coregonid populations in northern Fennoscandia (Østbye et al. 2006; Saisa et al. 2008; Præbel et al. in press). Moreover, support for L. Sinettjärvi being the source population for the invasion was found in the combined approach of a neighbor-joining tree and Bayesian clustering, where genetic homogeneity was shown for L. Sinettjärvi and the Inari-Pasvik watercourse populations whereas L. Kelujärvi formed its own cluster. Bayesian clustering and neighbor-joining trees have successfully been used in other studies for inference of routes and sources of biological invasions (see review by Estoup and Guillemaud 2010), and supportive results, as seen herein, provide strong evidence for the revealed pattern. L. Sinettjärvi and L. Inari also shared a higher number of private alleles than L. Kelujärvi and L. Inari, which provide additional support for L. Sinettjärvi being the founders, as time since introduction is too short for many new private alleles to accumulate. Addressing the number of shared private alleles between pairs of populations has been used in several other studies to identify the origin of invasions and migrations (e.g., Szpiech et al. 2008; Bell and Matocq 2011). It is also worth considering that the similarity in genetic variation of L. Sinettjärvi and L. Inari provide additional support for this origin, as biological invasions very seldom are associated with an increase in genetic variation in the introduced population compared to the source population (Nei et al. 1975; Dlugosch and Parker 2008b). Thus, taken together, there is compelling evidence that the second introduction of vendace in 1964–1966 from L. Sinettjärvi represents the source of the vendace invasion and establishment in the Inari-Pasvik watercourse.

Founder effects in the initial Inari-Pasvik vendace invasion

Multiple introductions have been shown to retain genetic diversity (Kolbe et al. 2004; Facon et al. 2008; Geller et al. 2010) and thus increase the likelihood of successful colonization and demographic expansion of the invader (e.g., Dlugosch and Parker 2008b; Shine 2012). In the present study the successful source population for the invasion was identified as the one that was introduced several times in 1964–1966 to a small lake upstream L. Inari. The population that apparently did not succeed in establishing was introduced to L. Inari by escapees from a local hatchery (Mutenia and Salonen 1992). This implies that the propagule pressure for this source likely has been smaller and/or that the population was presumably less adapted for the watercourse compared to the repeated introductions from L. Sinettjärvi. In fact, the results show that the L. Inari population has similar allelic richness and heterozygosity to the source population.

We did not identify loss of genetic variation via genetic bottlenecks for the introduced L. Inari population, which have been shown to be the case for other biological invasions (e.g., Tsutsui et al. 2000; Henry et al. 2009; Ayres et al. 2010). Similar effective population size estimates in the L. Sinettjärvi and Inari-Pasvik populations also suggested little genetic drift. Thus, the initial founding event of vendace in the Inari-Pasvik watercourse may be associated with no or limited founder effects. However, we show that the source and the invasive populations are genetically different, which suggest that genetic drift actually has played a role in the establishment of the colonists, since time from introduction is too short for mutations to have accumulated across the microsatellite loci in high enough frequencies to have effect on the result.

Dispersal pattern and genetic structure of the secondary expansion

The results revealed that the populations resulting from the secondary expansion downstream from L. Inari have diverged genetically from the introduced population in a stepping stone pattern. Such a dispersal pattern maintains genetic variation and limits divergence compared to other expansion patterns (e.g., Ibrahim et al. 1996; Reusch et al. 2010; Tonione et al. 2011), due to the allelic patchiness and likely subsequent gene flow among demes. The populations within the secondary expansion have become genetically distinguishable from the L. Inari population in about 18 years/9 generations (maximum 35 years/17 generations if the Pasvik populations were founded already in 1973 at the time of first observation in L. Inari, but this scenario appears highly unlikely from the observed occurrences of vendace in the watercourse; see Amundsen et al. 1999). This is remarkable, especially because the high genetic variation and large consensus and effective population size of the colonist population effectively should counteract genetic divergence. In marine fish species, such as herring (*Clupea harengus*), capelin (*Mallotus villosus*), and Atlantic cod (*Gadus morhua*), it is well known that large population sizes with high genetic variation and no obvious barriers to gene flow effectively hamper the build up of reproductive isolation (Knutsen et al. 2003; Mariani et al. 2005; Præbel et al. 2008; but see review by DeWoody and Avise 2000). The major difference and the most likely explanation for the rapid divergence observed among the vendace populations within the Inari-Pasvik watercourse therefore appears to be the presence of several hydroelectric dam constructions between the L. Inari population and the two downstream populations from the secondary expansion (i.e., L. Skrukkebukta). These dams may effectively hamper downstream

gene flow and promote the build up of reproductive isolation during the expansion.

Adaptive and evolutionary changes in the invader – perspectives

Identifying the source population and reconstructing the routes of biological invasions are crucial for handling and managing invasive species, as well as for gaining knowledge of the ecology behind successful colonization events. However, invasions also offer the possibility to study evolutionary processes such as natural selection and life history changes as they unfold. For example, invaders may evolve in response to an altered selection regime compared to their native range (Facon et al. 2006; Sax et al. 2007; Bacigalupe 2009; Shine 2012). The Inari-Pasvik watercourse is 300–380 km north of L. Sinettjärvi, but at similar altitude. Thus, the summer is shorter and colder, and the ice-covered period longer. Especially, temperature is an important driver for adaptive changes as it will affect, for example, egg incubation time, growth, physiological processes (Q10), and general individual fitness (e.g., Mooney and Hobbs 2000). In addition, r-strategists are often favored during biological invasions (Lodge 1993; Facon et al. 2006), and our earlier study suggests that these invasive populations in the secondary expansion areas have speeded up their life history via growth and age at sexual maturity (Amundsen et al. 2012). These changes happened within a decade and the system therefore apparently represents an excellent example of rapid adaptive evolution. Future studies of this system may therefore gain valuable insights by identifying adaptive phenotypic traits (transcriptome) and the genetic basis for local adaptation (e.g., using phenotypic QTLs) (see e.g., Bernatchez et al. 2010). Moreover, insights would be gained from investigating variation at genes and genomic blocks related to important life history parameters such as growth, metabolism, and disease resistance, to infer adaptive changes or patterns of plasticity among the Sinettjärvi-Inari-Pasvik populations.

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Conflict of Interest

None declared.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1. Summary statistics for each locus and population in the source and invader vendace (*Coregonus albula*) populations from northern Europe.

Table S2. Geographical distances and expansion times of vendace populations used in the partial Mantel test.

Table S3. Estimates of genetic differentiation (F_{ST})

between the five populations of vendace in northern Europe.

Figure S1. Summary of results from STRUCTURE for inferring K populations among the tested vendace populations when omitting three loci that may be associated with null alleles.

Figure S2. Summary of results from STRUCTURE for inferring K populations among the tested vendace populations.