Early germination to seed set in *Heracleum persicum*. Photos: Dilli P. Rijal
Reconstructing the invasion history of *Heracleum persicum* (Apiaceae) into Europe

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Abstract

Sparse, incomplete and inappropriate historical records of invasive species often hamper invasive species management interventions. Population genetic analyses of invaders might provide a suitable context for the identification of their source populations and possible introduction routes. Here, we describe the population genetics of *Heracleum persicum* Desf. ex Fisch and trace its route of introduction into Europe. Microsatellite markers revealed a significantly higher genetic diversity of *H. persicum* in its native range, and the loss of diversity in the introduced range may be attributed to a recent genetic bottleneck. Bayesian cluster analysis on regional levels identified three and two genetic clusters in the native and the introduced ranges, respectively. A global structure analysis revealed two worldwide distinct genetic groups: one primarily in Iran and Denmark, the other primarily in Norway. There were also varying degrees of admixture in England, Sweden, Finland and Latvia. Approximate Bayesian computation indicated two independent introductions of *H. persicum* from Iran to Europe: the first one in Denmark and the second one in England. Finland was subsequently colonized by English populations. In contrast to the contemporary hypothesis of English origin of Norwegian populations, we found Finland to be a more likely source for Norwegian populations, a scenario supported by higher estimated historical migration from Finland to Norway. Genetic diversity *per se* is not a primary determinant of invasiveness in *H. persicum*. Our results indicate that, due to either pre-adaptations or rapid local adaptations, introduced populations may have acquired invasiveness after subsequent introductions, once a suitable environment was encountered.

Keywords: approximate Bayesian computation, biodiversity, genetic variation, giant hogweeds, invasive alien species, population genetics

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Introduction

Invasive alien species affect biodiversity at all organizational levels from genes to ecosystems (Vitousek & Walker 1989; Vilà et al. 2011), and cause significant damage to the environment and economy (Pimentel 2011). Interspecies hybridization between the invasive and native species is considered a major cause for loss of native genetic distinctness (Rhymer & Simberloff 1996; Lockwood et al. 2013). Moreover, invasive alien species can change entire ecosystems by altering fire regimes (Pemberton & Ferriter 1998; Brooks et al. 2004; Watt et al. 2009; Simberloff 2013), hydrology (Zavaleta 2000), fauna of decomposers (Bedano et al. 2014) and nutrient pools (Vitousek et al. 1987; Wang et al. 2015).
Invasive alien species are considered one of the major threats to global biodiversity (CBD 2001; Genovesi et al. 2013). Besides, considerable concern in understanding biological invasion, management, control, and eradication of invasive species remains challenging due to sparse, incomplete and inappropriate historical records (Estoup & Guillemaud 2010). Due to this lack of historical information, many invasive species remain unnoticed until their populations explode. However, indirect methods based on molecular genetic markers have proved effective in bridging such gaps between invasion history and management by providing insight into the complex history of biological invasions (Lombaert et al. 2014).

Information about population genetics, introduction history and identification of source populations are crucial in understanding the invasion process (Cristescu 2015). The genetic diversity of a species indicates its evolutionary potential to adapt to a novel environment (Sakai et al. 2001). This may be especially important for exotic invasive species as they have to adapt and survive to novel environments. Genetic diversity of introduced populations largely depends on the number of founders and the number of introductions from the genetically differentiated (native) source populations (Kolbe et al. 2004; Lavergne & Molofsky 2007; Ward et al. 2008; Simberloff 2009). Genetically diverse populations may have higher establishment success if they contain genetic variants more suited to the new environment, thereby posing greater invasion risk (Lee 2002; Forsman 2014; Bock et al. 2015). Although introduced invasive species suffer from genetic bottlenecks, they often overcome adverse effects of population reduction by genetic admixture via multiple introductions from the native range (Kolbe et al. 2004) and/or other successful introduced populations (invasive bridgehead effect, Lombaert et al. 2010; Benazzo et al. 2015). Given that multiple introductions and genetic admixture may enhance invasibility (Kolbe et al. 2004; Roman & Darling 2007; Marrs et al. 2008; Ward et al. 2008), the number of introductions may indicate risk of further regional spread of a species. Better understanding of the genetic diversity of introduced populations and vital source populations along with the number of introductions may be used to prevent further introductions and/or spread of invasive species by design monitoring and quarantine strategies targeting the source area and the important vectors (Estoup & Guillemaud 2010). Thus, genetic diversity of invasive populations can be used as a risk assessment tool.

The change in effective sizes and ranges of natural populations in the past leave signatures in their genetics (Cornuet et al. 2010), and this historical signature can be inferred by examining genetic variation among populations (Lawton-Rauh 2008). For example, genetic differentiation among populations is considered a product of limited dispersal and gradual genetic drift. As a result, genetic similarity becomes correlated with geographical distance (isolation by distance, Wright 1943). Introduction route of a species can be inferred using molecular data in several ways, including assessing similarity among genetic clusters (Pritchard et al. 2000; Bensard et al. 2014; Yu et al. 2014), assigning individuals to source populations (Ranula & Mountain 1997; Paetkau et al. 2004), quantifying gene flow between isolated populations (Nielsen & Wakeley 2001) and comparing plausible migration scenarios using simulation approaches (Beaumont et al. 2002; Cornuet et al. 2010; Bensard et al. 2014). Invasive vascular plants constitute about 53% of the invasive species of Europe, and 49% of these plants are of non-European origin (Pysek et al. 2009). Anthropogenic pressure is a main driver of European plant invasion, and a strong positive correlation is found between human population density and alien richness (Marini et al. 2012). Most alien plant species have deliberately been introduced into Europe, ornaments in particular (Lambdon et al. 2008). Among the many terrestrial invasive plant species, a group of large hogweeds commonly known as ‘giant hogweeds’ are posing threats to public health and biodiversity in different parts of Europe (Nielsen et al. 2005; EPPO 2009). Giant hogweeds (sensu Nielsen et al. 2005) include three invasive species of Heracleum (Apiaceae) in Europe (i.e. H. mantegazzianum, H. persicum and H. sosnowskyi). The first two species were famous garden plants during the 19th century in Europe, and the latter was introduced into northwest Russia as a forage crop at the end of the 1940s (Nielsen et al. 2005; EPPO 2009; Alm 2013). Within <2 centuries of introduction, giant hogweeds became some of the most prominent invasive species in northern Europe. They possess some typical features of invasive species, for example early and fast growth, high stature, huge biomass production, extensive cover and abundant seed production. In addition, H. persicum is perennial and highly clonal, which is not the case for other two giant hogweeds. It has successfully adapted to new environmental conditions, from hot summers of Persia, with ‘short’ days, to the much cooler conditions and perpetual daylight in parts of its introduced range at 51–71° northern latitude. An invasive species possessing all the characteristics of the ‘ideal-weed’ (Baker 1965) rarely exists in nature; however, H. persicum seems to exhibit most of the necessary characteristics (van Kleunen et al. 2015). Thus, H. persicum represents a model to provide broader understanding of the evolution of invasiveness, especially the paradoxical role of population bottlenecks, genetic diversity of the source populations, and introduction history.

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The source and introduction route of H. persicum in Europe are unclear. Hypotheses concerning introduction routes are based on historical accounts and limited observational data (Estoup & Guillemaud 2010). The first seed record of H. persicum in Europe comes from the seed list of Royal Botanic Garden Kew from 1819 (Pysek et al. 2010). Historical records show that an English man planted seeds in northern Norway in 1836 (Christy 1837; Fröberg 2010; Alm 2013); however, it is unclear whether he brought seeds from naturally growing English populations or from other sources. Meanwhile, the absence of naturalized populations of H. persicum in the UK (Sell & Murrell 2009; Stace 2010) is surprising, as the species has proved highly invasive elsewhere in NW Europe. In addition, the taxonomy of the giant hogweeds has been a subject of controversy (Jahodová et al. 2007; Fröberg 2010; Alm 2013), and a variety of ill-defined Latin names have been used for Scandinavian plants, including H. giganteum, H. lacinium, and H. panaceae. Heracleum persicum may be hiding in historical accounts due to misinterpretation as H. mantegazzianum. Under such circumstances, population genetics of H. persicum may serve as a promising alternative to resolve not only introduction pathways, but also illuminate the complex invasion history (Estoup & Guillemaud 2010; Brouat et al. 2014).

Even though H. persicum is highly invasive in the introduced range, we assume that it suffered a loss of genetic diversity due to population bottlenecks during the initial introduction. To test whether introduced populations are genetically depauperate, we compared the genetic diversity of native and introduced populations. Introduced populations often overcome the effects of genetic bottlenecks due to multiple introductions or genetic admixture, and we considered the number of introductions as an indicator of propagule pressure that may enhance establishment success of H. persicum. We evaluated whether introduced populations were formed by multiple introductions and if there has been admixture between introduced populations. To aid management interventions, we identified respective source populations of the introduced invasive populations and tested whether genetic diversity per se was inherently linked with invasiveness. By tracing the routes of introduction, we evaluated whether H. persicum followed the route indicated by historical accounts when invading Europe.

Material and methods

Study species

The enigmatic, invasive Heracleum species found in northern Scandinavia has been identified as Heracleum persicum based on genetic similarity with Iranian species (Jahodová et al. 2007), which is also supported by morphological investigations (Fröberg 2010). Although earlier studies (Nielsen et al. 2005; EPPO 2009; Fröberg 2010) stated that H. persicum was native to Iran and Turkey, Ahmad (2014) has recently reported it as a new species in Iraq, at a single station close to the Iranian border. Similarly, H. persicum is narrowly distributed in southeast Turkey (GE Ana
tolia) (Ahmad 2014; Arslan et al. 2015) in an area bordering northwest Iran. However, it is widely distributed in north, west, northeast and central Iran (Rechinger 1987; Ahmad 2014). It was introduced to Denmark, England, Finland, Latvia, Norway, Sweden and Iceland (Fröberg 2010; Wasowicz et al. 2013). The plant is polycarpic and generally attains a height of 2.5 m and sometimes reaches up to 3 m (Fröberg 2010; Alm 2013). Seed germination requires stratification at 2–4 °C for two months and flowering starts after the third year post germination. Temporal variation in flower maturation promotes outcrossing. Male flowers in the primary umbel mature earlier than female flowers. In the secondary umbels, flowering occurs after seeds are set in the primary umbels, and female flowers are generally abortive (Often & Graff 1994; Fröberg 2010). Reproduction primarily occurs through seeds; however, clonal reproduction is also common in disturbed habitats where seed reproduction fails. The plant sap is phototoxic and induces photoprotective allergy when exposed to ultraviolet radiations (Nielsen et al. 2005; EPPO 2009). In the introduced range, H. persicum commonly grows at seashores, roadsides, abandoned farmlands, highly disturbed areas and seminatural habitats like forest clearings. The earliest European record of the species appeared in the seed list of Royal Botanic Gardens, Kew, London, in 1819 (Pysek et al. 2010). It has been recommended for regulation as a quarantine pest in Europe (EPPO 2009) and is blacklisted in Norway (Gederaas et al. 2012).

Plant material

Historical records of the species from the global biodiversity information facility (GBIF) (http://www.gbif.org/species/3628745), Norwegian Biodiversity Information Centre (http://www.biodiversity.no/), sampling locations reported by Jahodová et al. (2007) and the most recent data available for Norway (Fremstad & Elven 2006) were rigorously evaluated before starting the sampling (Fig. 1). Sampling was done throughout the species’ distribution range between 2012 and 2014 (Fig. 1), except Iraq and Iceland, for which the species has only recently been found (Wasowicz et al. 2013; Ahmad 2014), and Turkey, from where export of plant material is now prohibited. We collected four...
samples and one representative herbarium voucher from 5 different spots at 5–10-m intervals per population, and care was taken to avoid resampling from the same genet, resulting in 1–20 samples per population. All samples were dried on silica gel and photographed. A few populations collected during 2003–2004 were retrieved from the material of Jahodová et al. (2007) (see Table 1) and herbarium vouchers for those samples are

Fig. 1 Geographical locations of previous records (small circles) and genetic structure of sampled populations from native and introduced ranges of *Heracleum persicum*. Size of a pie chart reflects gene diversity (expected heterozygosity) of each population. Hatched and plain pie charts indicate proportion of genomes of each population assigned to Cluster I and Cluster II, respectively, as revealed by global structure analysis based on *K* = 2. Dates indicate the first seed and plantation record for England and Norway (bold), respectively, the first cultivation record for Tromsø (bold italic), and the earliest records of garden escapes for Scandinavia (normal). Arrow indicates inferred route of introduction of *H. persicum* into Europe based on approximate Bayesian computation analysis.
Table 1. Sampling details and genetic diversity indices for populations of Heracleum persicum. Populations with <4 samples (italicized) were not considered while calculating average diversity indices across country (bold).

<table>
<thead>
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<th>District/Region</th>
<th>Location</th>
<th>Latitude</th>
<th>Longitude</th>
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<th>Year</th>
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<th>P (%)</th>
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<td>Umeå</td>
<td>63.8237</td>
<td>20.2783</td>
<td></td>
<td>DPR</td>
<td>2013</td>
<td>2.0</td>
<td>40.00</td>
<td>0.28</td>
<td>1.44</td>
<td>1.39</td>
<td>NA</td>
<td>0.28</td>
<td>0.20</td>
<td>0.26</td>
<td>-0.43</td>
</tr>
</tbody>
</table>

Sample collectors: AP, Atehfeh Pirany; DPR, Dilli Prasad Rijal; GG, Gertrude Gavrilova; IGA, Inger Greve Alsos; LF, Lars Fröberg; MFA, Mohsen Falahati-Anbaran; OB, Olaf Booy; PU, Pertti Uotila; RS, Rouollah Sobhian; SJ, Sarka Jahodova; TA, Torbjorn Alm; TJ, Tina Jorgensen; N, number of samples; P (%), percentage of polymorphic loci; I, Shannon’s information index; N_A, average number of alleles over loci; N_E, effective number of alleles; A_R, allelic richness based on three samples; H_O, observed heterozygosity; H_E, expected heterozygosity; uH_E, unbiased expected heterozygosity; F_IS, inbreeding coefficient; NA, not applicable.
deposited with original collectors. The leaf samples, DNA extracts, and herbarium vouchers of all other samples are deposited at Tromsø Museum (TROM).

**DNA extraction and standardization**

DNA was extracted using a DNeasy Plant Mini Kit (Qiagen, Hilden, Germany) following manufacturer’s protocol. DNA concentration of each sample was measured by NanoDrop 2000 (Thermo Scientific, Waltham, MA, USA), and all the samples were normalized to 10 ng/µL for downstream analyses.

**Microsatellite genotyping**

We selected 25 microsatellite markers developed by Rijal et al. (2015) and two markers developed by Henry et al. (2008), the latter two accommodated in multiplex II and III of Rijal et al. (2015), to genotype microsatellites of *H. persicum*. Altogether, 578 samples of *H. persicum* were screened in three multiplexes as described by Rijal et al. (2015). The total volume of PCR was 6 µL, which consisted of 3 µL master mix and 0.5 µL RNA-free water (Type-it Microsatellite PCR Kit; Qiagen), 1 µL primer mix and 1.5 µL template DNA. The thermal cycling conditions of each multiplex PCR were as follows: initial denaturation at 95 °C for 10 min followed by 10 cycles of 95 °C for 30 s, 60–50 °C of touch down PCR for 1 min with 1 °C decrease per cycle, and 72 °C for 45 s; 25 cycles of 95 °C for 30 s, 50 °C for 1 min, 72 °C for 45 s; and a final extension of 60 °C for 15 min. A mixture of 2 µL of 1:20 diluted PCR product, 7.8 µL of HiDi Formamide and 0.2 µL of LIZ 600 (Applied Biosystems, Foster City, CA, USA) was denatured at 95 °C for 5 min, and electrophoresis was performed on 3130xL genetic analyzer (Applied Biosystems). Samples that had poor amplification or failed during fragment analysis were re-analysed. Any samples with poor chromatogram, after re-analysis, were discarded from genotyping. The genotyping error rate (Bonin et al. 2004) was estimated by replicating 96 samples for 7 loci from multiplex III.

**Data analysis**

The fragments were further analysed in GENEIOUS version 6.1.6 (Biomatters Ltd, New Zealand) following 3rd-order least squares method implemented in microsatellite plugin for allele calling. Due to stutter band in locus Hp_25, allele calling became problematic in some of the populations. The locus Hp_05 was polymorphic for only one sample from Denmark. Thus, we discarded these loci from further analyses. Similarly, three samples were discarded from the further analysis due to poor chromatograms. PGDSPIDER version 2.0.5.0 (Lischer & Excoffier 2012), MICROSATELLITE TOOLS (Park 2001) and GENEALEX version 6.5 (Peakall & Smouse 2012) were used as data conversion tools, and the latter two were also used to check errors in genotypic data. Genotypic error rate was estimated by taking the ratio of mistyped genotypes to the total observed genotypes during the replication (the per-genotype error rate) whereas the ratio of miss-called allele to the total number of observed allele in the replication was considered as the per-allele error rate (Morin et al. 2009).

**Hardy–Weinberg equilibrium and linkage disequilibrium**

The test of Hardy–Weinberg equilibrium (HWE) and linkage disequilibrium (LD) was performed in GENEPOP version 4.3 (Raymond & Rousset 1995; Rousset 2008) with 10 000 dememorization and in 1000 batches with 10 000 iterations per batch. We also performed a HWE jackknife test (Morin et al. 2009) using package ‘strataG’ (Archer 2014) in r version 3.1.2 (Team 2014) to detect the influential samples in populations. We reran the HWE test to evaluate the impact of influential samples on HWE by omitting samples with unusually large odds ratio (>99% of the rest of the distribution) as suggested by Morin et al. (2009).

**Molecular diversity and genetic differentiation.**

The percentage of polymorphic loci (P%), Shannon’s information index (I), unbiased expected heterozygosity (UHE), average (N_A) and effective (N_E) number of alleles, observed (H_O) and expected heterozygosity (H_E), inbreeding coefficient (F_IS), and frequencies of private alleles were calculated for populations with ≥4 samples, that is 38 populations and 25 loci. All the analyses were performed in GENEALEX version 6.5 (Peakall & Smouse 2012).

Allelic richness (A_R) was calculated to account for the possible bias due to difference in population size. The pairwise population genetic differentiation (F_ST) was calculated and tested for significance based on 1000 permutation without assuming HWE. Both analyses were performed in FSTAT version 2.9.3.2 (Goudet 1995). FSTAT is sensitive to missing loci and produces error while calculating A_R and does not provide P-values for F_ST. The locus Hp_30 was not present in Danish populations; loci Hp_07, Hp_10 and Hp_24 were missing in Latvia, and in Gryllefjord locus Hp_23 was present in two individuals. Thus, we included populations with nine or more samples (30 populations) and excluded the aforementioned loci, that is, 20 loci included, while calculating A_R and F_ST. Null alleles overestimate population differentiation by reducing within-population genetic diversity. The frequency of null allele was estimated following expectation maximization (EM)
algorithm (Dempster et al. 1977) as implemented in FREENA (Chapuis & Estoup 2007). The global $F_{ST}$ was calculated with and without correction for null allele, using FREENA with 1000 bootstrap resampling over loci, to evaluate the impact of null alleles in estimation of genetic differentiation.

Native and introduced populations were not equally represented in this study due to unequal sampling. Thus, when comparing diversity estimates between native and introduced ranges we used Welch two sample t-test, which corrects the problem of unequal sampling by incorporating variance in the analysis and adjusting the degrees of freedom (Ruxton 2006). The tests were performed in R version 3.1.2 (Team 2014).

**Genetic bottleneck.** To assess the effects of population bottlenecks in *H. persicum*, tests of heterozygosity excess and deficiency, were performed in BOTTLENECK version 1.2.02 (Piry et al. 1999), using all available mutation models, with 1000 iterations. Infinite allele model (IAM) overestimates, whereas stepwise mutation model (SMM) underestimates the bottleneck signature (Cornuet & Luikart 1996). Two-phase mutation model (TPM) is one of the complex but realistic mutational models that also includes the possibility of non-stepwise mutations to SMM (Selkoe & Toonen 2006). Thus, a TPM was used with 70% proportion of SMM along with 30% variance for TPM. To get an overview, results based on all mutation models were evaluated by applying Wilcoxon’s test as it is the most powerful method when <20 polymorphic loci are considered (Cornuet & Luikart 1996; Piry et al. 1999). We also used mode shift test available in BOTTLENECK version 1.2.02 (Piry et al. 1999) to explore the recent bottleneck-induced distortion in the allele frequency (Luikart et al. 1998; Awad et al. 2014). The signature of subsequent population expansion after the bottleneck was tested with k and g tests (Reich & Goldstein 1998) using an excel macro KGTESTS (Bilgin 2007). Populations with ≥4 samples, that is 38 populations and 20 loci, were included in both of the analyses.

**Population genetic structure.** All 25 loci and 575 samples from 50 populations (Table 1) were assessed for genetic relationship by principal coordinate analysis (PCoA) in GENALEX version 6.5 (Peakall & Smouse 2012). The number of genetic clusters in *H. persicum* was estimated in STRUCTURE version 2.3.4 (Pritchard et al. 2000). The genetic structures of native and introduced populations were first evaluated separately. Altogether, 25 loci and 548 samples from 38 populations (with ≥4 samples) from native and introduced ranges were included in a global analysis. To detect the most likely native sources of the introduced populations, Denmark, England and Finland were analysed separately as well as jointly with native populations. To identify the likely sources of Norwegian populations, they were analysed separately with English and Finnish populations as well as in combination with all others. The analysis was performed on the Lifeportal computing platform (https://lifeportal.uio.no/) with initial burnin period of 200 000 followed by 250 000 Markov Chain Monte Carlo steps. The independent allele frequency and admixture model was assumed when performing Bayesian clustering analyses. The expected number of clusters (K) was set to 1–10 with 10 iterations for each K. The structure output was further processed in STRUCTURE HARVESTER (Earl & vonHoldt 2012). The best K was selected based on the Evanno et al. (2005) as implemented in STRUCTURE HARVESTER (Earl & vonHoldt 2012). Finally, summation of the individual file for different runs from STRUCTURE was performed in CLUMPACK (Kopelman et al. 2015).

**Colonization routes.** To trace the most likely introduction route of *H. persicum* in Europe, we tested four competing hypotheses by implementing approximate Bayesian computation (ABC) approach in DRY-ABC version 2.0.4 (Cornuet et al. 2014). Sweden and Latvia consisted of only 8 and 6 multilocus genotypes without missing loci, respectively, and their genetic structures were similar to England and Finland. The addition of less informative populations not only increases the number and complexity of the ABC scenarios, but also poses challenges in the result interpretation (Estoup et al. 2012). Thus, Latvia and Sweden were excluded from the ABC analysis, and 20 random multilocus genotypes without missing genotypes were selected each from England, Finland, Iran and Norway, and 19 from Denmark. The theoretical rationale for such regional sampling is provided in Stensien et al. (2011).

Testing historical scenarios within the ABC framework is inherently a post hoc analysis, and the hypotheses (historical scenarios) are generally based on the available historical information and genetic population structures (Estoup et al. 2012; Lombaert et al. 2014). Our hypotheses were also based on historical records and we used genetic evidence to test those hypotheses. Most of the introduced alleles (nearly 78%) were in a subset of Iranian alleles, and private alleles of the introduced range were seemingly recently mutated from alleles introduced from Iran (Table S3). Thus, we tested the following scenarios (Fig. 2) by considering Iranian populations as the native source of the introduced populations: (i) scenario 1 was based on the historical account which assumes that *H. persicum* was first introduced from Iran to England and then to Norway, and finally to Denmark and Finland from Norway; (ii) scenario 2 assumed serial introductions from Iran to Denmark to
England to Finland to Norway; (iii) scenario 3 assumed two independent introductions from England to Denmark and from Denmark to Finland, while Finland acted as source for Norway; and (iv) scenario 4 hypothesized two independent introductions from Iran to Denmark and England. The Finnish population was assumed to have originated in England and acted as source for Norwegian populations.

The priors in the ABC analysis were defined based on the available information and later adjusted according to the results of initial runs. The effective population size of the native range (Iran) and introduced ranges were considered as \( N_1: 10-2000 \) and \( N_2: 10-200 \), respectively. Due to high abundance of \( H. \) persicum in Norway, but low genetic diversity, different ABC runs were performed assuming effective size of Norwegian population equal to Iran as well as less than or equal to other introduced populations. Invasive species suffer through an initial bottleneck as only few individuals invade the new area (Sakai et al. 2001). \( H. \) persicum produces hermaphrodite flowers and like most of the members of Apiaceae, the species is considered to be self-compatible (Perglová et al. 2007). On this basis, we assume that even a single plant of \( H. \) persicum can produce seeds. Thus, we arbitrarily specified population size during bottleneck (\( N_1b \)) as 1–100. A variation of 30–100 years in the lag phase of invasive weeds has been reported (Aikio et al. 2010). If we assume the upper limit as the lag phase for \( H. \) persicum and a generation time of 3–6 years, then bottleneck duration may also vary from 17 to 33 generations. In general, defining narrow bottleneck duration prior reduces the accuracy of scenario identification (Guillemaud et al. 2010). Thus, we defined a wide period, that is 2–100 generations as the bottleneck duration (\( db \)). The species was present in Europe as early as 1819, which gives an estimate of 32–65 generations if we assume 3–6 years as the generation time of \( H. \) persicum. To cover the uncertainties in the divergence time we chose to use widely divergent time priors. Thus, the time since divergence of the recent to the oldest clades was considered as \( 2–100, 2–200, 2–300 \) and \( 2–400 \) generations ago and defined as \( t_1, t_2, t_3 \) and \( t_4 \), respectively. All the microsatellite loci were included in a single group and assumed to follow the identical mutation model with minimum mutation rate of \( 10^{-6} \) to maximum \( 10^{-2} \) per generation as reported for plant microsatellites (Udupa & Baum 2001; McConnell et al. 2007). The reference table was generated by \( 8 \times 10^6 \) randomizations, twice the number considered optimal by the program (Cornuet et al. 2014). We compared the posterior probabilities of competing scenarios based on the logistic regression of the raw and the linear discriminant analysis (LDA) transformed summary statistics (Estoup et al. 2012; Lombaert et al. 2014). We used \( 4 \times 10^6 \) simulated data sets while performing logistic regression on LDA transformed summary statistics. The type I and II error rates were used to discriminate the most plausible scenario. Type I error was the proportion of the number of times other scenarios have the highest posterior
probability than the scenario under consideration. Type II error rate was based on the scenario II which had the largest type II error rate (as suggested by Estoup et al. 2012) and calculated as the proportion of the number of times the scenario under consideration has the highest posterior probability in scenario II.

Migration rates. To quantify the demographic parameters, especially migration rate between Norway and Finland, we used isolation with migration analysis in IMa software, which allows subsequent migration between two lineages being split from an ancestral population (Nielsen & Wakeley 2001; Hey & Nielsen 2004, 2007). The isolation with migration analysis was performed setting the upper limit of the prior distribution of population mutation parameter as 1 for both Norway and Finland and 10 for the ancestral population. The upper migration priors for both lineages were set to 250. The divergence time prior for two lineages was set to 0.5. Burn-in period was set as 10 000 and genealogy was saved each hour. Metropolis coupling was implemented with 20 chains and two geometric heating terms, that is 0.8 and 0.9. Average mutation rate of microsatellite loci was considered as $10^{-5}$ (Udupa & Baum 2001; McConnell et al. 2007). Three replicates of isolation with migration analyses were performed with identical settings until 50 million MCMC steps had been generated after burn-in.

Results

Genotypic error

Four samples had a replicate with poor chromatograms and were removed from downstream analyses. The absolute difference between loci varied from 0.07 to 1.03 base pairs (bp) with mean ($\pm$SE) of 0.26 ($\pm$0.06) bp based on two replicates of 92 samples. We observed a per-genotype error of 2.2%, which was slightly higher than the per-allele error rate of 1.5%.

Hardy–Weinberg equilibrium and linkage disequilibrium

Out of 950 population-locus combinations, 37 departed from HWE after Bonferroni correction (about 4%, Table S1). Most of the combinations (29) deviating from HWE were confined to three loci – Hp_13, Hp_14 and Hp_20 – and the remaining eight deviations were distributed among populations, occurring no more than twice per population and locus (Table S1). Jackknife analysis produced odd ratios for loci Hp_14 and Hp_20, indicating that these two loci had a comparatively large impact on tests for deviations from HWE (result not shown). Removal of 18 samples with $\geq$1.2 odd ratio did not change the overall HWE result (result not shown). The test of genotypic disequilibrium was significant for two loci pairs (Hp_27 × Hp_30 and HMNSSR_132B × HMNSSR_206) after Bonferroni correction (Table S2).

Molecular diversity and genetic differentiation

The average percentage of polymorphic loci was lowest for Norway (52.1%) and highest for Sweden (86.4%) (Table 1). Out of 205 alleles recorded, 163 were common and 25 and 17 were private to the native and the introduced populations, respectively. There were 48 and 35 alleles private to native and introduced ranges, respectively (Table S3). The Latvian population did not contain any private alleles. The Shannon’s information index, allelic richness, expected and unbiased expected heterozygosities were lowest in Norway and highest in Iran (Table 1). The average number of alleles ranged from 1.72 (Latvia) to 3.34 (Iran). Minimum and maximum values of the observed heterozygosity were found for Norway and England, respectively. Similarly, the inbreeding coefficient ranged from −0.24 (England) to 0.11 (Iran). Locus-wise diversity statistics for native and invaded ranges are provided in Table S4.

Out of 435 comparisons, $F_{ST}$ values of 295 population pairs were significant after Bonferroni correction (Table S5). One population from Iran (Mazandara) was not significantly differentiated from any native or introduced populations (nonsignificant pairwise $F_{ST}$). Three populations from Norway (Kvaløyvegen of Tromso, Hammerfest and Nesna) were not significantly differentiated from most of the native and introduced populations. The mean ($\pm$SE) country-wise $F_{ST}$ (averaged over population) was lowest between England and Sweden, that is 0.267 ($\pm$0.006), and highest between Norway and Denmark, that is 0.552 ($\pm$0.005) (Table 2). The average ($\pm$SE) frequency of null allele per locus varied from 0 ± 0 to 0.140 ± 0 (Table S6). There was a strong positive correlation between number of alleles and frequency of null allele, and only five loci had $>0.05$ null allele frequency (Fig. S1, Supporting information). The average ($\pm$SE) frequency of null alleles per population ranged from 0.001 ± 0 to 0.137 ± 0.023 (Table S6). The genetic differentiation between native and introduced ranges remained nonsignificant, when $F_{ST}$ was estimated by including and excluding null alleles (result not shown).

The percentage of polymorphic loci, Shannon’s information index, average numbers of alleles, effective number of alleles, private alleles, allelic richness; observed, expected (gene diversity) and unbiased expected heterozygosities, as well as inbreeding coefficients were significantly higher in the native range than
in the introduced range (Table 3). The loss of genetic diversity ranged from 16 to 49% in the introduced range, and on average nearly 42% of the gene diversity \( (H_E) \), Table 3) was lost by the introduced populations compared to the native populations. The average frequency of null alleles was significantly higher in native compared to introduced range. The fixation index, \( F_{ST} \), was lower in the native compared to the introduced range, but the difference was marginal and nonsignificant (Table 3).

**Genetic bottleneck**

The tests of heterozygosity excess was significant after Bonferroni correction for one native and seven introduced populations when infinite allele model was considered (Table S7). However, the numbers were reduced to four and three introduced populations when two-phase and stepwise mutation models were assumed, respectively. Neither heterozygosity excess nor deficiency was observed in one native and twelve introduced populations. Similarly, mode of the allele frequency was shifted in 79% of the populations. About 67% native and 81% of the introduced populations showed mode shifts in the allele frequency distributions indicating recent bottlenecks (Table S7).

The within-locus \( k \) tests were significant for five introduced populations, indicating a signal of population expansion (Table S7). The interlocus \( g \) test was not very informative, as there were no clear trends between \( g \) ratios and significant \( k \) values (Table S7).

**Population genetic structure**

Ordination of microsatellites revealed that the Iranian, Danish and Norwegian populations of *Heracleum persicum* were distinct from each other. Populations from England, Finland, Latvia and Sweden appeared in between the former populations in the ordination plot (Fig. 3). Most of the variation (22.9%) in ordination plot was explained by the first axis while the second axis explained 6.6% of the variation. Finland consisted of highly variable samples scattering across most of the length of the first axis (Fig. 3).

There were three and two distinct genetic clusters in the native and the introduced ranges of *H. persicum*, respectively (Fig. 4). The two genetic clusters remained consistent when native populations were analysed with introduced populations from each country or in combinations (Fig. S2, Supporting information). Based on the rate of change of the likelihood distribution and the delta \( K \) value (Fig. 4C), two genetic clusters were detected for *H. persicum* in a global analysis (Figs 1 and 4D). More than 90% of the genomes of Norwegian samples were assigned to cluster I (hatched cluster in Figs 1 and 4D,F). However, more than 90% of the genomes of Iranian and Danish samples were assigned to cluster II (plain cluster in Figs 1 and 4D,F). Samples from England, Finland, Latvia and Sweden shared a higher

---

**Table 2** The country-wise \( F_{ST} \) values averaged over populations of *Heracleum persicum*. Standard errors are given in the parentheses.

<table>
<thead>
<tr>
<th>Country</th>
<th>Iran</th>
<th>Denmark</th>
<th>England</th>
<th>Finland</th>
<th>Latvia</th>
<th>Norway</th>
<th>Sweden</th>
</tr>
</thead>
<tbody>
<tr>
<td>( F_{ST} )</td>
<td>0.253 (0.023)</td>
<td>0.388 (0.015)</td>
<td>0.037 (0.000)</td>
<td>0.385 (0.014)</td>
<td>0.336 (0.010)</td>
<td>0.082 (0.000)</td>
<td>0.409 (0.019)</td>
</tr>
<tr>
<td>( F_{ST} )</td>
<td>0.407 (0.019)</td>
<td>0.452 (0.009)</td>
<td>0.306 (0.003)</td>
<td>0.354 (0.025)</td>
<td>0.000 (0.000)</td>
<td>0.503 (0.006)</td>
<td>0.552 (0.005)</td>
</tr>
</tbody>
</table>

---

**Table 3** Comparison of overall genetic diversity statistics between native and introduced populations of *Heracleum persicum*.

<table>
<thead>
<tr>
<th>Estimates</th>
<th>Native</th>
<th>Introduced</th>
<th>( t )</th>
<th>d.f.</th>
<th>( P )-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>( P ) (%)</td>
<td>85.50</td>
<td>59.81</td>
<td>4.82</td>
<td>15.78</td>
<td>0.000</td>
</tr>
<tr>
<td>( I )</td>
<td>0.80</td>
<td>0.40</td>
<td>5.66</td>
<td>6.60</td>
<td>0.001</td>
</tr>
<tr>
<td>( N_A )</td>
<td>3.40</td>
<td>1.88</td>
<td>4.62</td>
<td>5.50</td>
<td>0.005</td>
</tr>
<tr>
<td>( N_E )</td>
<td>2.21</td>
<td>1.50</td>
<td>5.04</td>
<td>5.74</td>
<td>0.003</td>
</tr>
<tr>
<td>( A_R )</td>
<td>2.16</td>
<td>1.61</td>
<td>5.00</td>
<td>6.88</td>
<td>0.002</td>
</tr>
<tr>
<td>( H_O )</td>
<td>0.38</td>
<td>0.30</td>
<td>2.43</td>
<td>8.58</td>
<td>0.039</td>
</tr>
<tr>
<td>( H_E )</td>
<td>0.43</td>
<td>0.25</td>
<td>5.86</td>
<td>8.20</td>
<td>0.000</td>
</tr>
<tr>
<td>( uH_E )</td>
<td>0.45</td>
<td>0.27</td>
<td>5.88</td>
<td>8.43</td>
<td>0.000</td>
</tr>
<tr>
<td>( F_{IS} )</td>
<td>0.11</td>
<td>–0.14</td>
<td>3.95</td>
<td>10.88</td>
<td>0.002</td>
</tr>
<tr>
<td>( P_A )</td>
<td>4.17</td>
<td>1.89</td>
<td>3.07</td>
<td>7.89</td>
<td>0.016</td>
</tr>
<tr>
<td>( F_{ST} )</td>
<td>0.25</td>
<td>0.30</td>
<td>–1.94</td>
<td>19.77</td>
<td>0.066</td>
</tr>
<tr>
<td>Null allele</td>
<td>0.07</td>
<td>0.03</td>
<td>3.11</td>
<td>5.72</td>
<td>0.022</td>
</tr>
</tbody>
</table>

Nonsignificant \( P \)-value is in bold.

\( P \) (%), percentage of polymorphic loci; \( I \), Shannon’s information index; \( N_A \), average number of alleles over loci; \( N_E \), effective number of alleles; \( A_R \), allelic richness based on three samples; \( H_O \), observed heterozygosity; \( H_E \), expected heterozygosity; \( uH_E \), unbiased expected heterozygosity; \( F_{IS} \), inbreeding coefficient; \( P_A \), number of private alleles; \( F_{ST} \), fixation index.
proportion of both clusters. Assignment graphs of higher K values (2–4) for native, introduced, native-Denmark, native-England, global analyses and Norway are provided as supporting information (Fig. S2, Supporting information).

Colonization routes

The pre-evaluation of the scenarios suggested that priors were satisfactory delimited as the simulated data surrounded observed data in the ordination plot. There were no differences in the overall scenario discrimination.
patterns when the effective population size of Norway varied. The third scenario, which assumed two independent introductions from England to Denmark and Finland as well as another introduction to Norway from Finland, appeared more plausible than other scenarios when raw summary statistics were used. The posterior probability of scenario III was slightly higher in both the direct and logistic methods (average posterior probabilities 0.390 ± 0.010 and 0.648 ± 0.014, respectively) (Fig. S3 and Table S8, Supporting information). However, the highest posterior probability (0.651 ± 0.004) was observed for the fourth scenario, which assumed multiple introductions to Denmark and England from Iran, when LDA transformed summary statistics were used. The type I and II error rates were 3.0 and 1.9 times higher for the scenario III compared to the scenario IV, respectively, when using raw summary statistics (Table 4). The LDA transformed summary statistics produced 5.8 and 0.9 times higher type I and II error rates, respectively, for the scenario III compared to the scenario IV. The observed data of the scenario IV was more properly surrounded by the posteriors than the scenario III (Fig. S4, Supporting information), which further indicated that the fourth scenario was more likely than others.

The effective population sizes of Iran and Denmark/England/Finland/Norway under scenario IV were estimated to 1250 and 132, respectively (median of England/Finland/Norway under scenario IV were estimated to 1250 and 132, respectively (median of $N_1$ and $N_2$, Table 5). The result indicated that the Danish and the English lineages of $H. persicum$ were derived from Iran about 218 and 139 generations ago respectively (median of $t_4$ and $t_3$, Table 5). However, the Finnish and the Norwegian lineages were split from their respective common ancestors about 75 and 57 generations ago, respectively (median of $t_2$ and $t_1$, Table 5). The medians of the biases were found within the range of −0.046 to 0.839 for $t_1$ and $db$, respectively (Table S9).

**Table 4** Type I and II error rates for scenarios 3 and 4 (see Fig. 2 for the details) based on the logistic regression with raw (from $8 \times 10^8$ simulated data) and LDA transformed (from $4 \times 10^8$ simulated data) summary statistics

<table>
<thead>
<tr>
<th>Errors</th>
<th>Summary statistics</th>
<th>Type I</th>
<th>Type II</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Raw</td>
<td>LDA transformed</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.43</td>
<td>0.44</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.11</td>
<td>0.06</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3.0</td>
<td>5.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.26</td>
<td>0.25</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.09</td>
<td>0.14</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.9</td>
<td>0.9</td>
</tr>
</tbody>
</table>

LDA, linear discriminant analysis.

**Migration rate**

Exact mutation rates of Heracleum microsatellites have not been reported. When minimum ($4.4 \times 10^{-4}$) and maximum ($1.4 \times 10^{-3}$) mutation rate estimates from ABC analysis (Table 5, 25 and 97.5% quintiles) were used, population divergence time ($\tau/\mu$) varied from 24 to 75 generations for highest to lowest mutation rates. Average divergence time of Norwegian and Finnish lineages estimated by isolation with migration model was nearly 50 generations, which was approximately similar to the ABC estimates. The IM model suggested a higher rate of migration from Finland to Norway than vice versa (Table 6 and Fig. 5).

**Discussion**

We found significantly lower percentages of polymorphic loci, allelic richness and private alleles in the introduced range of Heracleum persicum compared to its native range. In addition, a significant loss of genetic diversity, as revealed by reduced expected heterozygosity and effective number of alleles, was also observed in the introduced range. Heterozygosity excess, an indicator of a genetic bottleneck, was observed in a few introduced populations.

**Genetic diversity, population differentiation and bottleneck**

Several monomorphic loci, lower genetic diversity, shifts in allele frequency and bottleneck signatures detected in the introduced range indicate that the introduced populations were established by few founders (Cornuet & Luikart 1996; Luikart et al. 1998; Piry et al. 1999; Sakai et al. 2001). Meanwhile, tests of recent population expansion was significant for five Norwegian populations growing south of Tromsø. Spread of H. persicum south of Tromsø is considered as a more recent event in Norway (Alm 2013). Successful invaders are expected to experience frequent bottlenecks without dramatic changes in genetic variation (Dlugosch et al. 2015). Thus, detection of bottleneck signature and population expansion characterizes a general process of initial establishment and colonization of H. persicum as it is spreading to new locations (Alm 2013; Wasowicz et al. 2013). Some of the earliest records of H. persicum in Norway come from Hammerfest, Honningsvåg, Talvik and Tromsø (see Fig. 1) (Alm 2013 & references therein), and none of them showed signatures of bottlenecks. Thus, evidence of bottlenecks is more common in the most recent populations, which agrees with general principles of the currently employed test that expect detection of bottleneck signatures for relatively
recently bottlenecked populations (2N_e-4Ne generations in the past) (Cornuet & Luikart 1996; Piry et al. 1999).

The inbreeding coefficients were significantly lower for introduced populations, indicating a genetic bottleneck. Inbreeding depression depends on several factors including life history stages and population history (Husband & Schemske 1996). In general, due to fewer individuals, mating between close relatives (biparental inbreeding) is nearly unavoidable in smaller populations, which could force species towards the verge of extinction as a consequence of inbreeding depression and loss of alleles (Newman & Pilson 1997; Frankham & Ralls 1998). Thus, one would expect severe inbreeding in introduced species, as they are generally founded by few individuals, which in turn may reduce fitness. Surprisingly, inbreeding coefficients were either close to zero (an indication of perfect outcrossing) or negative (an indication of heterozygote excess) for introduced populations of *H. persicum*. Inbreeding can be avoided and outcrossing promoted through protandry in Apiaceae, a feature that has been reported for *Heracleum mantegazzianum* (Perglová et al. 2007). Inbreeding coefficients close to zero for several native and introduced populations indicate that the phenomenon is pervasive in both ranges. Negative inbreeding coefficients, on the other hand, have been frequently reported for the introduced populations of invasive species (Walker et al. 2003; Henry et al. 2009; Hagenblad et al. 2015). Thus, it could perhaps be viewed as a phenomenon linked with reduction in population size during expansion of the invasive species. Populations which showed relatively more negative inbreeding coefficients were those that predominantly had bottleneck signatures under IAM (Table 1 and Table S7). Thus, populations exhibiting a significant heterozygosity excess or negative inbreeding coefficient might have experienced a recent genetic bottleneck (Cornuet & Luikart 1996).

### Table 5

ABC results of historical parameters estimated from 2008 pseudo-observed data sets simulated under scenario III (see Fig. 2) for *Heracleum persicum*. Mean, median, mode as well as 2.5, 5, 95 and 97.5% quintiles of estimated values are provided. N1 and N2, current effective population size of Iran and Norway, and Denmark, England and Finland, respectively; db, duration of bottleneck; N1b, population size during bottleneck; t1, t2, t3 and t4 time since divergence of the youngest to the oldest lineages (see Fig. 2 and text for details).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Mean</th>
<th>Median</th>
<th>Mode</th>
<th>q25</th>
<th>q50</th>
<th>q95</th>
<th>q97.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>N1</td>
<td>1250</td>
<td>1250</td>
<td>1250</td>
<td>528</td>
<td>637</td>
<td>1880</td>
<td>1940</td>
</tr>
<tr>
<td>N2</td>
<td>130</td>
<td>132</td>
<td>136</td>
<td>58</td>
<td>70</td>
<td>186</td>
<td>193</td>
</tr>
<tr>
<td>db</td>
<td>29</td>
<td>22</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>81</td>
<td>89</td>
</tr>
<tr>
<td>N1b</td>
<td>62</td>
<td>64</td>
<td>69</td>
<td>17</td>
<td>24</td>
<td>95</td>
<td>97</td>
</tr>
<tr>
<td>t1</td>
<td>56</td>
<td>57</td>
<td>54</td>
<td>19</td>
<td>25</td>
<td>87</td>
<td>92</td>
</tr>
<tr>
<td>t2</td>
<td>79</td>
<td>75</td>
<td>66</td>
<td>28</td>
<td>34</td>
<td>139</td>
<td>157</td>
</tr>
<tr>
<td>t3</td>
<td>144</td>
<td>139</td>
<td>142</td>
<td>54</td>
<td>65</td>
<td>242</td>
<td>261</td>
</tr>
<tr>
<td>t4</td>
<td>222</td>
<td>218</td>
<td>215</td>
<td>83</td>
<td>99</td>
<td>362</td>
<td>379</td>
</tr>
<tr>
<td>Mutation rate</td>
<td>0.00075</td>
<td>0.0007</td>
<td>0.0006</td>
<td>0.00044</td>
<td>0.00047</td>
<td>0.00122</td>
<td>0.0014</td>
</tr>
</tbody>
</table>

### Table 6

Maximum-likelihood estimates (MLE) along with the 95% highest posterior density (HPD) intervals for divergence time (τ = t1 where t is the generation since divergence) of Norwegian (N) and Finnish (F) lineages of *Heracleum persicum*. Estimates of ancestral (θA), Norwegian (θN) and Finnish (θF) population size as well as migration rate to Norway (mF > N) and to Finland (mN > F) are provided.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>95% HPD low</th>
<th>MLE high point</th>
<th>95% HPD high</th>
</tr>
</thead>
<tbody>
<tr>
<td>τ</td>
<td>0.015</td>
<td>0.033</td>
<td>0.474</td>
</tr>
<tr>
<td>θA</td>
<td>0.003</td>
<td>0.003</td>
<td>0.037</td>
</tr>
<tr>
<td>θN</td>
<td>0.003</td>
<td>0.003</td>
<td>0.037</td>
</tr>
<tr>
<td>θF</td>
<td>6.038</td>
<td>0.836</td>
<td>498.340</td>
</tr>
<tr>
<td>mF &gt; N</td>
<td>203.423</td>
<td>331.607</td>
<td>965.098</td>
</tr>
<tr>
<td>mN &gt; F</td>
<td>18.708</td>
<td>48.542</td>
<td>237.292</td>
</tr>
<tr>
<td>mF &gt; N</td>
<td>10.208</td>
<td>10.458</td>
<td>156.625</td>
</tr>
</tbody>
</table>

Fig. 5 Log-scaled marginal densities of migration rate of *Heracleum persicum* from Norway to Finland (dashed line) and Finland to Norway (solid line) estimated by IM analysis.

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In general, introduced populations are genetically less diverse than native populations (Barrett & Kohn 1991; Sakai et al. 2001; Lavergne & Molofsky 2007) and this is also the case for introduced and native populations of *H. persicum*. This pattern is expected when only a fraction of the genetic diversity of the native population is introduced during initial colonization (Barrett & Kohn 1991). In addition, introduced populations generally suffer from population bottlenecks often for a longer period of time, which also reduces the genetic diversity (Allendorf & Lundquist 2003). However, Dlugosch et al. (2015) argue that invaders often retain significant amount of genetic variation if the founding populations are large enough to overcome the demographic constraints. In a closely related species, *H. mantegazzianum*, Walker et al. (2003) found a large genetic differentiation among populations at different river catchments in the introduced range and credited the observed variation to several independent introductions and relatively large initial founder populations. Niinikoski & Korpelainen (2015) found high genetic differentiation and a modest level of genetic variation in the introduced Finnish populations of *H. mantegazzianum*. It should be noted that both studies had no comparison with the native range and thus the differentiation is relative. Similarly, while comparing genetics of giant hogweeds, Jahodová et al. (2007) found high overall genetic variability in the invaded ranges and concluded that the invasive populations were not affected by genetic bottlenecks. In contrast, by comparing native and introduced populations of *H. mantegazzianum*, Henry et al. (2009) found a significant reduction in the genetic diversity in the introduced range and concluded that a founder event might have occurred. In extreme cases, some of the Norwegian invasive populations of *H. persicum* have lost >65% of the genetic diversity compared to native populations (Nesna & Gryllefjord, Table 1); otherwise, on average 16–35% of the genetic diversity was lost in the other introduced regions. Although nearly 50% of the genetic diversity is lost by the Norwegian populations compared to native populations (average H_E, Table 1), *H. persicum* is most abundant and vigorous in Norway compared to other introduced areas. Although neutral genetic markers may be poorly correlated with quantitative traits (Merilä & Crnokrak 2001; Reed & Frankham 2001; McKay & Latta 2002), a low level of genetic diversity does not seem to limit the invasiveness in giant hogweeds. Genetic diversity *per se* appears less important in determining the invasiveness of *H. persicum* in the introduced range. Genetics of invasive species, thus, represents a paradox in terms of the role of genetic diversity in adaptability (Simberloff 2013; Edelaar et al. 2015).

**Route of introduction**

We found higher population structuring within the native range as indicated by three distinct genetic clusters. However, two genetic clusters were consistent when some of the initially established introduced populations (Danish and English) were analysed separately or in combination with native populations, and populations from north-central Iran appeared more likely to be the sources of these introduced populations (Fig. S2C and D, Supporting information). A global Bayesian cluster analysis and ordination plot revealed two pure and one admixed genetic structures for introduced populations of *H. persicum* (Figs 1 and 4D,F). Denmark and Norway were clustered separately with distinct genetic structures, whereas England, Finland, Latvia and Sweden showed admixed genetic structure. Based on this result, we inferred that the Danish and all introduced genotypes (except the Norwegian) originated from two independent introductions from the native range, and the Norwegian genotypes originated from one of the introduced populations composed of mixed genotypes.

Although we could not include samples from Turkey and Iraq, genetic diversity, structure analyses and the *post hoc* ABC analysis indicated Iran as the source area for the European *H. persicum*. Nearly 78% of the introduced alleles were subset of the Iranian alleles and the remaining 22% private alleles were seemingly recent deviants of the Iranian alleles (1–4 mutational steps, Table S3). Although our six populations covered the major geographical distribution of the species in Iran (see Fig. 1), relatively higher genetic differentiation among Iranian populations (Figs 4A and S2, Supporting information) indicates that inclusion of more populations from Iran would have encompassed most of the introduced private alleles. Nevertheless, the apparent similarity in the allelic composition between Iran and the introduced range of *H. persicum* is unlikely to be a chance effect alone. The narrow distribution of *H. persicum* in Turkey, as well as its morphological mismatch with the Scandinavian specimens (Øvstedal 1987), makes it less likely to assume Turkey (and even more so, Iraq, with only a single, recently discovered station 400 m from the Iranian border) as sources of the European *H. persicum*, although we cannot exclude this as those populations were not sampled. The wide distribution of *H. persicum* in Iran as well as its morphological and genetic similarity with the European specimens (Jahodová et al. 2007; Fröberg 2010) indicate Iran as the more likely source of the European *H. persicum*.

Our findings do not corroborate the contemporary hypothesis that assumes an English population of *H. persicum* as the source of Norwegian population and all other European populations as descendant of the
latter (Nielsen et al. 2005; Jahodová et al. 2007; EPPO 2009). In an earlier study, Jahodová et al. (2007) con-
cluded that, as the Danish population appeared com-
pletely different from other introduced populations but
more similar to Iran, multiple introductions from Iran
might be responsible for invasion of H. persicum in Nor-
dic countries. Structure analysis revealed that the Dan-
ish populations are more genetically similar to the
Iranian than to the other introduced populations. As
introduced populations tend to be more genetically sim-
lar to the source population(s) than to each other (Bond
et al. 2002), our data indicate that the introduced popu-
lations were founded by more than one independent
introduction from Iran.

In the ABC analyses, the LDA transformed summary
statistics provided the highest support for the scenario
IV that assumed two independent introductions to Den-
mark and England from the native source, and the sub-
sequent spread in other parts from England. Although
direct summary statistics provided the highest support
for the scenario III, we considered scenario IV as the
most likely scenario based on LDA transformed sum-
mary statistics. LDA reduces the number of dimensions,
which decreases the number of explanatory variables
and maximizes the differences among the scenarios,
thereby improving the accuracy of the ABC approxima-
tion by avoiding correlations among explanatory vari-
ables (Estoup et al. 2012). In addition, scenario IV had
lower type I and II error rates compared to scenario III.
The ABC result was also supported by Bayesian cluster
analysis showing shared clustering between English,
Norwegian and Finnish but not Danish populations
(Figs 1 and 4D,F). The genetic variation of introduced
populations depends on the genetic diversity of the
source population, and a relative decrease (due to bot-
tleneck) or increase (due to multiple introductions and
admixture) in the diversity of the introduced population
is likely to happen (Edelaar et al. 2015). However, nei-
ther structure analysis nor genetic diversity patterns
indicate any genetic admixture in the introduced range.
Multiple introductions do not seem to have increased
genetic variation. Instead, the pattern of loss of the
genetic diversity in the introduced range closely resem-
bled the introduction events indicated by the ABC anal-
yses. For instance, Danish and English populations
most likely originated from the similar native source
from Mazandaran of central Iran close to the capital city
Tehran (see Fig. S2, Supporting information) and have
lost nearly 16% and 19% of the genetic diversity of the
source; Finnish populations lost 6% of the English
genetic variation; and Norway lost nearly 33% of the
Finnish genetic variation. Thus, genetic diversity pat-
terns of H. persicum appear to have been shaped largely
by diversity of the source and the introduction history.

Although ABC appears as a promising methodology
for inferring invasion scenarios, incorporating too many
populations exponentially decreases the probability of
accepting a simulation, a phenomenon known as the
‘curse of dimensionality’. It also increases the number
of scenarios and parameters to be tested (Beaumont
et al. 2002; Cornuet et al. 2010). We traced the invasion
history of H. persicum by ABC analysis and expected
managers to utilize this information to avoid further
introduction by isolating or eliminating small, intro-
duced populations from the important source popula-
tions. We still suggest caution while interpreting ABC
outcomes as our results were based on only four com-
peting scenarios (out of 120) possible introduction sce-
narios.

Nevertheless, IM analysis provided new insights into
the spread of H. persicum into Europe. As migration rate
was higher from Finland to Norway than the reverse, it
is quite likely that Norwegian populations were
founded by Finnish propagules. Though the first seed
record for H. persicum comes from Royal Botanic Gar-
dens, Kew, the first verified Nordic escape record
comes from Finland from 1871 (see Fig. 1) (Fröberg
2010). The first verified record of species in Denmark
dates back to 1888 and the first Norwegian record to
1899 (Fröberg 2010). In contrast, the Norwegian records
of H. persicum cultivation date back to the 1830s
(Christy 1837; Fröberg 2010; Alm 2013). One probable
explanation for this discrepancy may be the lack of his-
torical records of H. persicum in Finland. In Denmark,
past authors failed to realize that the introduced plants
could belong to several species, generally interpreting
both extant stands and the historical records as relating
to H. mantegazzianum (e.g. Brondegaard 1990). Bronde-
gaard (1979; p.307) cites anecdotal evidence of introduc-
tion of (presumed) H. mantegazzianum to Denmark in
the 1830s. The timing is probably more reliable than the
mode (as packing material for statues) and route (from
Italy) of transport. In the light of our molecular data,
early cultivations in Denmark are likely to have
included H. persicum.

In addition, historical records of workers’ movement
from Finland to Norway, especially in the area where
H. persicum was first recorded, further link Finnish and
Norwegian populations of H. persicum. The earliest doc-
umented introduction of a large Heracleum species to
northern Norway was made by a British traveller, W.
Christy, in 1836. He visited Kåfjord at Alta and Ham-
merfest, and distributed seeds from England at both
stations (Christy 1837). In 1835, Kåfjord was the largest
single settlement in the otherwise sparsely populated
county of Finnmark, due to the English-owned and run
copper mines. In 1840, the mines employed 651 work-
ers, with Finns constituting the largest ethnic group,
outnumbering Norwegians (Moberg 1968; Nielsen 1995). It is probable that seeds from northern Norway may have been transferred to Finland and vice versa. Thus, while genetic data confirms the historical record of link between Finland and Norway, the inferred direction of spread is opposite.

Extensive populations of *H. persicum* in Norway suggest that it might be one of the oldest European populations. However, if Norwegian populations were older than Finnish and Danish populations, and founded the latter two, we should expect to observe higher level of polymorphisms in Norway than in other places. Norwegian populations are composed of quite distinct genotypes (Figs 1, 3 and 4; Fig. S2, Supporting information) and genetically highly structured compared to other regions (highest average regional F_{ST}, Table 2), indicating limited dispersal. Reduced gene flow is a prerequisite for local adaptation (Lenormand 2002). Thus, despite the lowest genetic diversity, spatially extensive populations in Norway may be due to local adaptations or success of pre-adapted genotypes from Iranian temperate mountains. These genotypes may be favoured in cool northern Norwegian climate compared to other countries. From its present distribution in Norway, it is evident that *H. persicum* thrives in the humid coastal areas with mild winters and avoids the drier inland areas with their cold winters, which may also explain the general scarcity of records of naturalized plants in Sweden and Finland. Also, fewer ornamental plants are able to thrive in northern Norway than England and Denmark may have increased its popularity. The current genetic (dis)similarity among regional populations might be due to discrepancy in regional climate and local adaptation.

Management implications

The genetic diversity of *H. persicum* is comparatively lower in the introduced than in the native range. *Heracleum persicum*, however, is vigorous and highly invasive in the introduced range despite lower genetic diversity.

As it is now generally regarded as an obnoxious weed in Norway, we assume that the historical vector (i.e. frequent cultivation in gardens) responsible for the original introduction and dispersal of *H. persicum* is now obsolete, indicating no further risk of intentional introductions from the native sources (unless Iranian immigrants are tempted to cultivate it from fruits imported for culinary use). However, a successfully established invasive population may pose greater risk of spread than the native source as the former needs a single evolutionary shift to acquire invasiveness while the latter needs multiple changes along with independent evolution of traits to be invasive (Estoup & Guillemaud 2010; Lombaert et al. 2010). Further introduction and expansion of *H. persicum* are quite likely in Europe due to high frequency of cross-border travels and transportations. While tracing the route of the introduction of *H. persicum*, the English and the Finnish populations appeared as the important sources for founding introduced populations. We urge managers to pay special attention while formulating management interventions to avoid the possible second introduction from the respective sources. Otherwise, successive waves of introduction from similar sources may augment further invasions (Benazzo et al. 2015). In addition, population admixture due to multiple introductions is considered a stimulus for rapid evolutionary changes (Kolbe et al. 2004; Lavergne & Molofsky 2007; Facon et al. 2008; Dlugosch et al. 2015). Thus, it is important to emphasize that some populations in the introduced range of *H. persicum* (i.e. Denmark, England, Finland, and Sweden) still have higher genetic diversity and may contribute to increase genetic diversity of neighbouring populations, for example Norwegian populations, by multiple introductions.

In general, biological control agents are chosen from the native (source) range of the invasive species (Roderick & Navajas 2003). *Heterodera persica*, a cyst-forming nematode, has been reported to parasitize on *H. persicum* in Iran (Maafi et al. 2006). *Heterodera persica* may be considered as a candidate biocontrol agent in the introduced range of *H. persicum*; however, so far, there has been no effort to test the effectiveness of *H. persica* as biological control agent against *H. persicum*. Meanwhile, we suggest to carefully assess the pitfalls of biological control agents as it has received both negative and positive responses (Messing & Wright 2006; Seastedt 2015). Moreover, it is important to note that single agent from the native range adapted against certain genotypes of *H. persicum* may not be sufficient for biological control (Marrs et al. 2008) as there are two distinct and one admixed groups of *H. persicum* in Europe.

Most microsatellite markers used in this study are also polymorphic for other giant hogweeds, that is *H. mantegazzianum* and *H. sosnowskyi*, the native *H. sphondylium* which has been reported to hybridize with giant hogweeds (EPPO 2009), their invasive hybrids and some also for *Anthricus sylvestris* (Rijal et al. 2015). Hybridization can impede management interventions through creation of unique characteristics, for example production of novel chemicals, which in turn makes hybrids unrecognizable or unpalatable to specific herbivores or biological control agents (Schoonhoven et al. 2005; Williams et al. 2014). In general, hybridization appears a common phenomenon within the genus *Heracleum* (EPPO 2009). In particular, *H. persicum*
commonly hybridizes with *H. sphondylium*, producing fertile and vigorous hybrids. They have already shown their presence and effect in Scandinavia (Fröberg 2010; Alm 2013; Rijal *et al.* 2015), and may further pose management challenges due to enhanced invasive abilities in hybrids as a consequence of interspecies hybridization (Ellstrand & Schierenbeck 2000; Schierenbeck & Ellstrand 2009). Thus, population genetics of *H. persicum* may shed light on the genetic attributes of other giant hogweeds as well as their invasive attributes.

**Conclusions**

Even though the genetic data indicated at least two independent introductions of *H. persicum* to Europe, a clear genetic bottleneck was inferred, increasing with the stepwise introduction to more northern ranges within Europe. In contrast to the contemporary hypothesis of English origin of Norwegian populations, Finland appears as a more likely source for Norwegian populations of *H. persicum*. Despite the lowest level of genetic diversity, Norwegian populations are the most vigorous in the introduced range, suggesting no effect of bottlenecks on the invasiveness of *H. persicum*. Thus, genetic diversity *per se* does not seem to be an important determinant of invasiveness in *H. persicum*. Our result indicates that, due to either pre-adaptations or rapid local adaptation, introduced populations may acquire invasiveness after subsequent introductions when a suitable environment is encountered.

**Acknowledgements**

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**References**


Data accessibility

Geographical coordinates and sampling locations are provided in Table 1. DNA and primer sequences of 25 microsatellite markers used in this study are available as supplementary material in Rijal et al. (2015) at http://link.springer.com/article/10.1007/s11105-014-0841-y. Final microsatellite genotypes for 25 loci and 575 samples, input files and important analysis scripts are available on Dryad doi: http://dx.doi.org/10.5061/dryad.kg66r

Supporting information

Additional supporting information may be found in the online version of this article.

Fig. S1 Relationship between number of alleles per locus and null allele frequency in *Heracleum persicum*.

Fig. S2 The graphical output of structure analysis based on K = 2–4 for (A) Iran, (B) introduced range, (C) Iran-Denmark, (D) Iran-England, (E) global analysis, and (F) Norway. Based on the delta K value, the best K = 3 for Iran and 2 for all other analyses.

Fig. S3 Comparison of four scenarios for introduction history of *Heracleum persicum* based on (A) direct, and logistic regression with (B) raw and (C) LDA transformed summary statistics as implemented in DIYABC (see Fig. 2 and the text for details). X-axes, the number of simulated data closest to the observed; and y-axes, posterior probabilities.

Fig. S4 Principal component analysis of priors (open circle), posteriors (solid circle) and observed data (yellow solid circle) for (A) scenario III and (B) scenario IV (see Fig. 2 and the text for details).

Table S1 Exact test of Hardy-Weinberg equilibrium using a Markov chain with 10 000 demorization steps (1000 batches and 10 000 iterations per batch).

Table S2 Test of linkage disequilibrium using a Markov chain with 10 000 demorization steps (1000 batches and 10 000 iterations per batch).

Table S3 Number of private alleles in the native and the introduced ranges of *Heracleum persicum* along with their frequencies and nearest alleles.

Table S4 Locus wise genetic diversity statistics between native and introduced populations of *Heracleum persicum*.

Table S5 Pairwise population *F*<sub>ST</sub> values significantly different from each other after Bonferroni correction are indicated in bold.

Table S6 Estimation of frequency of null alleles in each population and locus by expectation maximization algorithm.

Table S7 Test of the signature of bottleneck following infinite allele model (IAM), two-phase model (TPM) and step-wise mutation model (SMM).

Table S8 Comparison of four historical scenarios of route of introduction of *Heracleum persicum* based on direct and logistic regression with raw and LDA transformed summary statistics as implemented in DIYABC.

Table S9 ABC results for mean relative biases of historical parameters for *Heracleum persicum* based on present data.