

# Population genetics of the common guillemot *Uria aalge* in the North Atlantic: geographic impact of oil spills

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**ABSTRACT:** The population genetic structure of a species can be an important conservation tool informing us about the potential for genetic loss and the capacity for species recovery. Depending on the nature of population subdivision, it can also provide a means for assessing the source population of dead or injured individuals of unknown geographic origin. This type of information can be particularly useful in instances of large-scale environmental accidents, such as oil spills. Following the wreck of the 'Erika' oil tanker in the Bay of Biscay in December 1999, more than 80 000 seabirds were washed ashore along the west coast of France. The most heavily affected species (80% of all birds) was the common guillemot *Uria aalge*, a widespread long-lived colonial seabird. In an attempt to evaluate the 'true' geographic extent of this accident, we carried out population genetic analyses using 6 microsatellite markers on samples from 22 breeding colonies in the North Atlantic and on individuals collected after the oil spill. A pattern of isolation by distance was detected among common guillemot populations, but populations were only weakly structured, even at large spatial scales. The low level of genetic differentiation between colonies prevented clear assignments of oiled birds to their population of origin using only the genetic information. The weak genetic structure suggests that little genetic variability was lost during the oil spill and implies a high potential for population recovery via dispersal. However, current gene flow among extant colonies may overestimate the capacity of locally extinct colonies to recover due to the behavioural processes involved in recruitment. Clearly, the mortality of tens of thousands of high-latitude seabirds due to oil pollution warrants ongoing scientific scrutiny and conservation effort.

**KEY WORDS:** Assignment tests · Dispersal · Microsatellites · Marine pollution · Common guillemot · *Uria aalge* · Colonial seabird

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## INTRODUCTION

Certain seabird species are particularly vulnerable to oil spills due to their ecology, behaviour and life-history traits (Piatt & Lensink 1989, Piatt et al. 1990, Wiens et al. 1996, Begg et al. 1997, Irons et al. 2000, Peterson et al. 2003). The colonial breeding habits of most seabirds, with colonies varying in size from tens to hun-

dreds of thousands of individuals, and their strong natal philopatry (tendency to return to breed where they were born) and breeding-site fidelity (Furness & Monaghan 1987, Harris et al. 1996a,b) could make populations particularly sensitive to local extinctions or to losses of large numbers of individuals. In addition, due to the high annual survival rate of adults and low annual fecundity (several species lay only 1 egg per

season), seabird populations can be highly sensitive to reductions in the survival of breeders (Wiens et al. 1984, Lebreton & Clobert 1991). Moreover, changes in the condition of the local habitat can also lead foraging individuals to move elsewhere, thereby influencing habitat occupancy and use (Wiens 1996). Depending on the time of the year when an oil spill occurs, breeding colonies closest to the spill site are not necessarily the most heavily affected. Birds from numerous geographic locations often gather together at sea during winter, a time when inclement weather may increase the frequency of oil spills, illegal discharges and shipwrecks. Such migratory groups can be particularly vulnerable. The relative importance of these different effects is difficult to assess due to the large scales often involved and the logistical constraints in monitoring the fate of birds at sea. In this context, determining the origins of seabirds caught in an oil spill is critical for any attempt to assess its potential impact (Piatt & Lensink 1989, Heubeck et al. 2003, Peterson et al. 2003), and population genetic approaches could provide valuable information.

Following the 'Erika' wreck in December 1999, more than 20 000 tonnes of oil were released into the Bay of Biscay and spread along the west coast of France (Le Moigne & Laubier 2004). This is an important wintering area for a large number of seabird species (Mead 1974, Brown 1985, Harris & Swann 2002), and more than 80 000 birds were found oiled along the French coast (Cadiou et al. 2004) Société pour l'Etude et la protection de la nature en Bretagne (Bretagne-Vivante-SEPNB) and Ligue pour la protection des oiseaux (LPO). Of these individuals, 80 % were common guillemots, *Uria aalge* (Family Alcidae), of unknown origin.

To investigate the geographic origins of birds affected by the 'Erika' oil spill, several approaches could be used. The first is based on the biometrical features of recovered birds (Hope Jones 1996, Stratford & Partridge 1996, Heubeck et al. 2003). For instance, the common guillemot is classified into 6 sub-species distributed along a north-south gradient in the North Atlantic. However, the actual description and distinctness of these sub-species is controversial (Bédard 1985, Brown 1985). Biometric methods are difficult to implement without a consensus on the morphological features to be used to characterize sub-species (Brown 1985). In addition, the spatial resolution of such a method can be imprecise (Cadiou et al. 2004). The second potential approach is based on ring recoveries (Baillie & Mead 1982, Harris & Swann 2002), which, while more reliable than biometry, also has some limitations. Typically, few oiled birds carry a ring, and as ringing has been intense in some areas but not in others, any inferences made from such techniques would likely be biased (Mead 1974, Harris & Swann 2002, Cadiou et al. 2004). A third approach to determine the geo-

graphical origin of affected individuals could be to use variation in the isotopic composition of sampled tissues, a method which reflects the natural variation of stable isotopes in the environment (Hobson 1999). Previous studies on seabirds have shown the potential of this technique to study migratory and wintering areas (Minami & Ogi 1997, Cherel et al. 2000), but investigations in this field are currently limited by a relatively poor understanding of marine biogeographic isotopic regions and the identification of small-scale isotopic gradients in food webs (Hobson 1999, Forero & Hobson 2003).

Here, we decided to use a fourth method that relies on highly polymorphic genetic markers to study the population structure of the common guillemot and to attempt to assign individuals to their population of origin. Assignment tests have been used to (1) identify individual dispersers (Paetkau et al. 1995, Rannala & Mountain 1997), (2) estimate rates of dispersal between populations (Favre et al. 1997), (3) reconstruct genetic lineages (Nielsen et al. 1997), and (4) determine the origin of individuals for conservation or wildlife management (Edwards et al. 2001, Manel et al. 2002, Maudet et al. 2002, Scribner et al. 2003). The principle of these tests is to compare the multilocus genotype of an individual to genotypes found in different sampled populations. Each individual can then be assigned to the population where it has the strongest probability of belonging (Paetkau et al. 1995, Waser & Strobeck 1998). The usefulness of this method depends on (1) the set of sampled populations and (2) the ability to characterize populations using the genotypes of individuals, which is higher if the populations are genetically structured (Cornuet et al. 1999). Studying population genetic structure can also provide information on natural levels of gene flow at different scales. This can be particularly useful for conservation as it gives us an idea of the risks of losing genetic variability and the capacity of local populations to recover via rescue effects and re-colonization following dramatic declines (Friesen 1997, Amos & Balmford 2001, Hedrick 2001).

Predictions concerning the genetic structuring of seabird populations are conflicting. Until the last decade, population studies on intensively monitored sites have shown high philopatry and breeding-site fidelity in several widespread species, such as the common guillemot (Tuck 1961, Cramp 1985, Hudson 1985, Harris et al. 1996a), which could lead to population differentiation. However, it is now well known that young auks may settle and start breeding away from their natal colony (e.g. Harris & Wanless 1991, Halley & Harris 1993, Lyngs 1993, Harris et al. 1996a, Olsson et al. 1999, Harris & Swann 2002). The effective dispersal of individuals between colonies could thus be strong enough to genetically homogenize populations. Weak

population genetic differentiation has been reported for several seabird species (e.g. Atlantic puffin *Fratercula arctica*: Moen 1991; Brünnich's guillemot *Uria lomvia*: Birt-Friesen et al. 1992; common guillemot: Moum et al. 1991, Friesen et al. 1996b, Moum & Arnason 2001; sooty tern *Sterna fuscata*: Avise et al. 2000; albatrosses, *Thalassache* spp.: Burg & Croxall 2001, Adélie penguin *Pygoscelis adeliae*: Roeder et al. 2001; marbled murrelet *Brachyramphus marmoratus*: Congdon et al. 2000; northern fulmar *Fulmarus glacialis*: Burg et al. 2003; but see great frigatebird *Fregata minor*: Dearborn et al. 2003). Most of these studies have used markers that typically reflect the phylogeographic history of the species rather than provide information on current gene flow (e.g., mitochondrial DNA markers), and a call for the use of more polymorphic markers, such as microsatellites, has been made (Moum & Arnason 2001). Microsatellites have become the marker of choice for many conservation studies because of their (1) expected neutrality, (2) abundance and broad distribution throughout the genome, and (3) relatively high polymorphism, even in bottlenecked populations and species with little polymorphism at other markers (e.g. Jarne & Lagoda 1996, Parker et al. 1998, Gaggiotti et al. 1999, Sunnucks 2000).

The aim of this study was to use a population-genetics approach as a general tool to investigate the geographic impact of marine pollution on seabird species. In particular, we considered the 'Erika' oil spill, and the most heavily affected species, the common guillemot. We evaluated the genetic structure of common guillemot populations in the North Atlantic using neutral, hypervariable molecular markers (microsatellites) and samples from a large number of colonies. As previous molecular studies on this species used more historical markers (Moum et al. 1991, Friesen et al. 1996b, Moum & Arnason 2001), this analysis provides new information on inter-colony gene flow of this abundant species in the North Atlantic. Using individuals sampled in different breeding colonies and ringed birds recovered following the oil spill, we then tested whether we could determine with certainty the geographical origin of oiled birds.

## MATERIALS AND METHODS

**The common guillemot.** The common guillemot breeds in the Northern Hemisphere between 40° and 70°. The North Atlantic breeding population is estimated at 3 million pairs, of which 85% are found along the eastern coast (Harris & Wanless 2004). Northern colonies can be large (tens of thousands of breeding pairs), but colonies situated at the southern limit of the range, e.g., in France, Spain and Portugal, are typically

small. Guillemots, like other auks, spend much time on the sea surface, making them particularly susceptible to oil pollution (Harris & Wanless 2004). They are long-lived birds, reaching sexual maturity at about 5 years of age (Harris et al. 2000). After overwintering at sea, they return in early spring to their breeding colony, typically on high, steep sea cliffs (Tuck 1961). They usually remain with the same mate between years and produce a maximum of 1 chick per year (Hudson 1985). Until recently, the individuals of this species were considered strongly philopatric and faithful to their breeding colony (Tuck 1961, Hudson 1985, Harris & Wanless 1995, Harris et al. 1996a,b). Nevertheless, it is now well known that young guillemots visiting other colonies may settle and start breeding away from their natal colony (Halley & Harris 1993, Lyngs 1993, Harris et al. 1996a).

**Sampling and genotyping.** A total of 678 common guillemot individuals were sampled from 22 colonies spread across the North Atlantic. Particular attention was given to the eastern North Atlantic (Britain and Ireland), the areas most likely to have been affected by the 'Erika' oil spill (Fig. 1, Table 1). As seabirds may form local kin groups within colonies (Friesen et al. 1996a), we attempted to sample in different areas within each colony. Samples from within each colony came from either breeding birds or chicks, to diminish the risks of sampling related or non-breeding individuals. In most colonies, blood samples were taken, although growing feathers and embryos were also used for DNA extractions (Table 1).

Following the oil spill, an effort was made to keep a representative sample of oiled birds. Among the large number of recovered guillemots, 1500 carcasses, gathered from several locations along the French coast, were frozen and used for DNA sampling. Samples of pectoral muscle from these individuals were collected and preserved in 95% ethanol. To evaluate the usefulness of the assignment method for determining the origins of these birds, we restricted our initial analysis to 48 ringed individuals for which the colony of origin was known (47 birds ringed in Britain, 1 in Germany).

DNA was extracted using a commercial kit (Perfect gDNA Blood Mini Isolation kit, Eppendorf). The same kit was used for solid tissue and feathers. All individuals were genotyped at 6 microsatellite loci. Three of these markers had been isolated previously for this species (Uaa5–8, Uaa23, Iburguchi et al. 2000; HC6, GenBank accession number: AY359959). Three others were developed for related species (K32 from black-legged kittiwake *Rissa tridactyla*, Tirard et al. 2002; Ulo22 and Ulo29 from Brünnich guillemot *Uria lomvia*, Iburguchi et al. 2000). For 3 loci (Uaa5–8, Ulo22 and Uaa23), polymerase chain reactions (PCR) were carried out as outlined in McCoy & Tirard (2000) using an

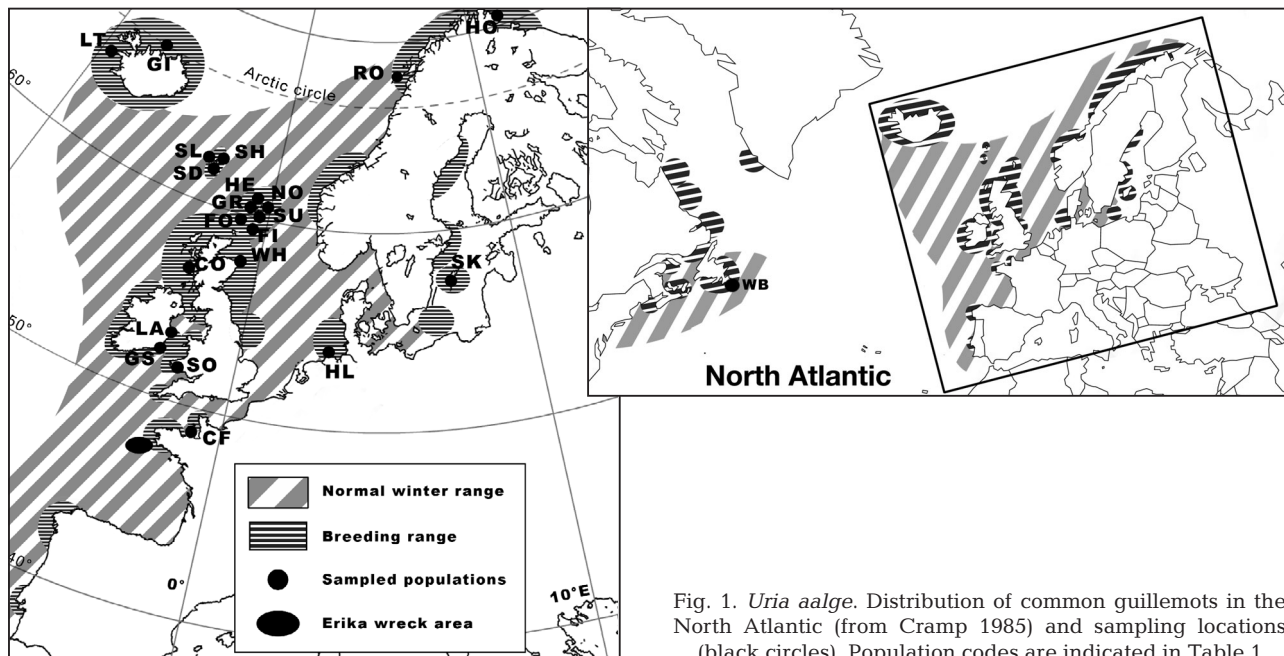


Fig. 1. *Uria aalge*. Distribution of common guillemots in the North Atlantic (from Cramp 1985) and sampling locations (black circles). Population codes are indicated in Table 1

annealing temperature of 55°C. PCR products were run on 6% polyacrylamide gels using size controls, and alleles were scored manually by 2 independent observers. The 3 other loci (Ulo29, Hc6, K32) were analysed using an automated sequencer (ABI Prism 310 Genetic Analyser, Applied Biosystems, Perkin-Elmer). For these markers, amplifications were performed in a 10 µl mixture containing approximately 50 ng of genomic DNA, 75 µM of each dNTP (10 mM

TrisHCl, 50 mM KCl and 1.5 mM MgCl<sub>2</sub>), 0.4 µM of each primer and 0.25 U Taq DNA polymerase (Qiagen). Primers were labelled with fluorescent dyes. Amplifications were performed in a GeneAmp PCR System 9700 (Applied Biosystem) with initial denaturation of 12 min at 94°C, 30 cycles of 15 s at 94°C, 15 s at 53°C and 30 s at 72°C, and a final elongation step of 10 min at 72°C. Alleles were scored via a labelled size standard (GENESCAN ROX 400HD) using the ABI

Table 1. *Uria aalge*. Origins and numbers of common guillemot samples

Colony	Latitude	Longitude	Number	Sample type	Individuals	Origin
Hornøya, Norway (HO)	70° 23' N	31° 09' E	24	Blood	Adults	T. Boulinier, K. McCoy
Røst, Norway (RO)	67° 28' N	11° 59' E	34	Blood	Adults	T. Anker Nilssen
Grimsey, Iceland (GI)	66° 33' N	18° 00' W	28	Blood	Adults	T. Boulinier, K. McCoy
Latrabjarg, Iceland (LT)	65° 29' N	24° 28' W	28	Blood	Adults	T. Boulinier, K. McCoy
Sandoy Liraberg, Faeroe isles (SL)	61° 54' N	06° 56' W	15	Embryos	Embryos	B. Olsen
Sandoy Honin, Faeroe isles (SH)	61° 52' N	06° 55' W	30	Embryos	Embryos	B. Olsen
Skuvoy Daer, Faeroe isles (SD)	61° 46' N	06° 49' W	33	Embryos	Embryos	B. Olsen
Hermaness, Scotland (HE)	60° 50' N	00° 55' W	30	Blood	Adults	T. Boulinier, K. McCoy
Grunay, Scotland (GR)	60° 39' N	01° 18' W	30	Blood	Adults	T. Boulinier, K. McCoy
Noss, Scotland (NO)	60° 09' N	01° 01' W	30	Blood	Adults	T. Boulinier, K. McCoy
Foula, Scotland (FO)	60° 08' N	02° 05' W	33	Blood	Adults	T. Boulinier, O. Duriez
Sumburgh Head, Scotland (SU)	59° 51' N	01° 16' W	31	Blood	Adults	T. Boulinier, K. McCoy
Fair Isle, Scotland (FI)	59° 32' N	01° 39' W	26	Blood	Adults	T. Boulinier, K. McCoy
Whinnyfold, Scotland (WH)	57° 23' N	01° 51' W	40	Blood	Adults	T. Boulinier, K. McCoy
Stora Karlso, Sweden (SK)	57° 17' N	17° 58' E	30	Feathers	Chicks	H. Osterblom
Helgoland, Germany (HL)	54° 12' N	07° 53' E	53	Feathers	Chicks	O. Hueppop
Colonsay, Scotland (CO)	56° 04' N	06° 13' W	32	Blood	Adults	T. Boulinier, K. McCoy
Lambay, Ireland (LA)	53° 29' N	06° 01' W	32	Feathers	Chicks	O. J. Merne
Great Saltee, Ireland (GS)	52° 07' N	06° 37' W	32	Feathers	Chicks	O. J. Merne
Skromer, Wales (SO)	51° 45' N	05° 17' W	30	Blood	Adults	T. Birkhead
Cap Fréhel, France (CF)	48° 25' N	02° 40' W	22	Blood	Chicks	T. Boulinier
Witless Bay, Canada (WB)	47° 16' N	52° 48' W	35	Blood	Adults	V. Friesen



Prism GENESCAN analysis software (v 2.1). For the samples from the 2 Icelandic colonies, all loci were analysed using the automated sequencer with the above protocol.

**Data analysis. Population genetic structure:** Fisher's exact probability tests, employing a Markov chain, were performed to test for departure from Hardy–Weinberg expectations for each population and locus, and for tests of linkage disequilibrium across populations (GENEPOP v3.3, Raymond & Rousset 1995). Significance levels were corrected for multiple tests (Rice 1989). Estimates of gene diversity (Nei 1987) and allelic richness in each population were computed using FSTAT (V2.9, Goudet 1995) and were compared among populations using a Kruskal-Wallis test (SAS Institute 1996). Population genetic structure was investigated using Wright's  $F$ -statistics (Weir & Cockerham 1984). The significance of  $F$ -statistic estimates was tested using 5000 permutations based on resampling alleles or genotypes either among individuals or among populations (FSTAT v2.9, Goudet 1995). Confidence intervals were obtained by bootstrapping over loci (Goudet 1995). To examine gene flow in common guillemots at different spatial scales, we tested for isolation by distance using the correlation between genetic distance, measured as  $F_{st}/(1 - F_{st})$ , and geographic distance, measured as  $\ln(\text{distance})$ , of population pairs (Rousset 1997). As common guillemots do not fly over land, geographic distances corresponded to the shortest distances by sea between 2 colonies (km). Correlations were tested for significance using a Mantel permutation procedure (Mantel 1967) associated with Spearman rank correlation coefficients (GENEPOP v3.3, Raymond & Rousset 1995).

**Assignment tests:** Assignment tests are only valid when populations are genetically structured (Cornuet et al. 1999, Manel et al. 2002, Maudet et al. 2002). To determine the reliability of the assignment tests for identifying the geographic origin of oiled birds, we first tested the method by attempting to assign the 678 individuals sampled in the 22 different colonies to their population of origin. As a second test, we used the same methods for assigning the 48 oiled ringed birds of known origin. We used the program GENECLASS (v4, Cornuet et al. 1999) which offers several methods for assigning individuals. We chose to use a likelihood-based method based on a Bayesian approach derived from the assignment method of Rannala & Mountain (1997) for computing the likelihood of a multilocus genotype occurring in a population. This method uses the allele frequencies of each population to compute the probability of occurrence of each multilocus genotype. We first used a direct assignment protocol to attribute each individual to the population to which its multilocus genotype had the highest probability of

belonging. Then, to estimate the statistical certainty of each individual assignment, we used a simulation exclusion method to compute the probability of excluding each candidate population (using 10 000 simulated individuals, significance threshold = 0.01, Bayesian estimation of frequencies, and the 'Leave one out' procedure, which excludes the tested individual from its population when estimating allele frequencies). For each candidate population, this method compares the likelihood of occurrence of an individual's genotype to the distribution of genotypes generated by Monte Carlo simulations. If the likelihood of the genotype is outside the tail of the distribution (significance threshold), the population is excluded as the population of origin of the individual.

## RESULTS

### Population genetic structure

No linkage disequilibrium was observed between any of the 6 loci. After correction for multiple tests, only 3 populations—Foula, Helgoland and Witless Bay—were in Hardy–Weinberg disequilibrium (Table 2). For Foula and Helgoland, disequilibrium was caused

Table 2. *Uria aalge*. Estimates of genetic variability (mean  $\pm$  SE) within common guillemot populations for 6 microsatellite loci. p-values refer to tests for deviations from Hardy–Weinberg equilibrium for each population. n refers to the number of individuals genotyped,  $r$  to allelic richness, and  $h$  to gene diversity. See Table 1 for colony abbreviations. Fis: Wright's inbreeding coefficient

Colony	n	$r$ (SE)	$h$ (SE)	Fis	p
HO	21	5.48 (1.05)	0.68 (0.07)	0.108	0.030
RO	34	5.69 (1.17)	0.70 (0.06)	0.007	0.581
GI	28	5.18 (0.91)	0.68 (0.06)	0.065	0.036
LT	28	5.09 (0.88)	0.67 (0.08)	0.131	0.145
SL	15	5.05 (1.09)	0.67 (0.07)	-0.051	0.052
SH	30	5.30 (0.96)	0.69 (0.06)	0.071	0.106
SD	33	5.09 (1.04)	0.64 (0.08)	0.000	0.941
HE	29	5.35 (0.88)	0.66 (0.05)	0.008	0.807
GR	28	5.32 (0.96)	0.67 (0.06)	-0.034	0.161
NO	30	5.75 (1.01)	0.69 (0.06)	0.101	0.053
FO	31	5.52 (0.98)	0.70 (0.05)	0.009	<0.00001
SU	31	5.26 (0.91)	0.68 (0.04)	0.053	0.560
FI	26	5.05 (0.88)	0.65 (0.06)	0.050	0.242
WH	40	5.54 (1.09)	0.69 (0.05)	0.018	0.700
SK	29	5.20 (0.97)	0.64 (0.06)	0.031	0.282
HL	49	5.55 (0.92)	0.68 (0.05)	0.107	<0.00001
CO	32	4.90 (0.87)	0.66 (0.06)	0.006	0.286
LA	27	5.26 (1.02)	0.69 (0.06)	0.021	0.284
GS	32	5.58 (1.01)	0.71 (0.05)	-0.016	0.894
SO	28	5.45 (1.01)	0.71 (0.06)	0.125	0.241
CF	22	5.65 (0.84)	0.70 (0.05)	0.048	0.558
WB	35	5.02 (1.24)	0.64 (0.10)	0.286	0.000

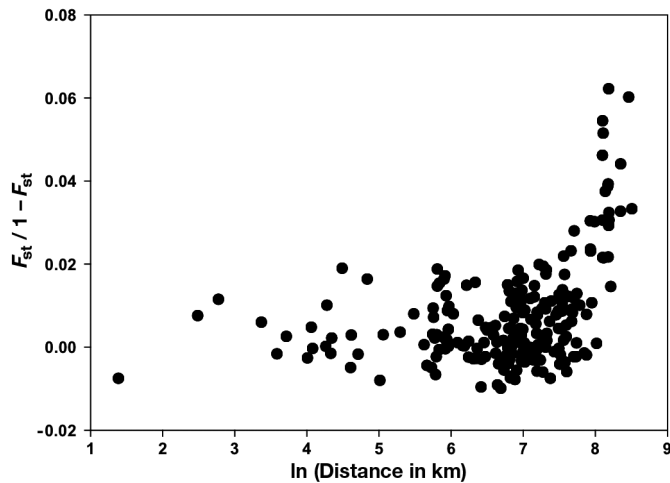


Fig. 2. *Uria aalge*. Pairwise genetic distance ( $F_{st}/1 - F_{st}$ ) vs. geographic distance ( $\ln[\text{distance}]$  by sea) between populations of common guillemot

by a single locus in each population (K32 and Uaa23 respectively). For Witless Bay, however, 3 loci (K32, Ulo22 and Hc6) were in disequilibrium. The problem at the K32 locus, which also showed slight heterozygote deficits in other populations, is likely due to the presence of a null allele (Callen et al. 1993), as this marker was developed for another species. We show results that include K32, but all tests were also carried out without this locus, to verify that results were unaffected by it. DNA samples from Helgoland were difficult to amplify and the disequilibrium at the locus Uaa23 could be due to the difficulties in reading gels. Finally, because more than 1 locus was affected for Witless Bay, sub-structuring within this population (i.e. the Wahlund effect, Hartl & Clark 1997) likely explains its deviation from Hardy–Weinberg equilibrium. Due to these 3 populations, alleles were not considered to be independent for testing the significance of population differentiation; permutation tests of  $F_{st}$  used the genotype as the randomization unit instead of the allele (Goudet 1995).

Among populations, the average allelic richness varied from 4.90 ( $\pm 0.87$ ) to 5.75 ( $\pm 1.01$ ), and gene diversity ranged from 0.64 ( $\pm 0.06$ ) to 0.71 ( $\pm 0.05$ ) (Table 2). However, neither parameter differed significantly among populations (Kruskal-Wallis tests;  $\chi^2_{21} = 1.86$ ,  $p = 1$  and  $\chi^2_{21} = 2.52$ ,  $p = 1$  respectively). Overall population genetic differentiation was weak (average  $F_{st} = 0.005$ ,  $p = 0.0014$ , 95% CI = 0.004 to 0.008). Results were similar when we excluded Witless Bay, the only population from the West Atlantic (average  $F_{st} = 0.004$ ,  $p = 0.0036$ , 95% CI = 0.002 to 0.007). In pairwise exact tests for population differentiation, only 3 population pairs were significantly differentiated after correction

for multiple tests: Witless Bay and Skuvoy, Witless Bay and Grunay, and Latrabjarg and Stora Karlsö (see Table 3). For each of these 3 population pairs, differentiation was explained by a single locus.

Genetic and geographic distances were correlated (Fig. 2;  $n = 231$ , Spearman's  $r = 0.39$ ,  $p < 0.0001$ ). This relationship seems to show a threshold effect at distances greater than 3000 km, distances that correspond to populations on either side of the North Atlantic. The same test performed without the most distant population (Witless Bay) resulted in a weaker, but still significant, correlation ( $n = 210$ , Spearman's  $r = 0.19$ ,  $p = 0.0051$ ). Thus, isolation by distance was principally explained by Witless Bay, the only population representing the western North Atlantic (Table 3).

### Assignment tests

One of the assumptions of assignment tests is that there is Hardy-Weinberg equilibrium within the population (Cornuet et al. 1999). We therefore excluded the 3 populations for which this condition was not valid (Foula, Helgoland and Witless Bay) from this analysis. When we used the direct assignment method, only 6% of the 563 birds sampled from breeding colonies were correctly assigned to the population from which they were sampled (Fig. 3), i.e. only 6% of the individuals

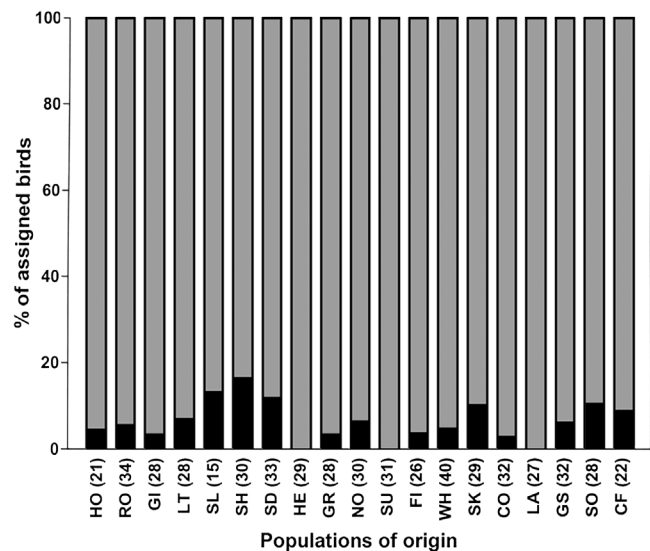


Fig. 3. *Uria aalge*. Direct assignments of the birds sampled from breeding colonies to populations. The black bars represent the percentage of individuals assigned to the population in which they were sampled, and the grey bars the percentage of birds assigned to another population. Note that we did not test samples from the populations which were in Hardy-Weinberg disequilibrium (Foula, Helgoland and Witless Bay). The numbers in brackets indicated the sample size of each population. See Table 1 for colony abbreviations

were genetically closer to their population of origin than to any other sampled population. The results obtained using the exclusion simulation method were similar in that most birds could be assigned to several populations. Indeed, for over 70% of the 563 individuals sampled in the breeding colonies, none of the 19 populations were excluded as the possible source. The same general results were obtained for the 48 ringed birds. Overall, the weak level of genetic structure prevented reliable use of assignment tests.

DISCUSSION

Our study showed that common guillemot populations in the North Atlantic are only weakly differentiated at microsatellite loci. This is surprising when we consider that this species shows strong natal philopatry and breeding-site fidelity, and it exhibits recognized morphological polymorphisms at the population level: North Atlantic common guillemots have been classified into several sub-species based on plumage and morphometric differences among colonies (Bédard 1985, Brown 1985), and a north–south cline in the proportion of the bridled morph of individuals has been reported (Birkhead 1984).

Because the performance of assignment tests is strongly influenced by the level of population differentiation (Cornuet et al. 1999, Manel et al. 2002, Maudet et al. 2002), the weak differentiation in common guillemots means that we were not able to determine the origin of the oiled birds with much certainty. Indeed, only 6% of sampled individuals could be directly assigned to their colony of origin and for more than 70% of them, none of the 19 populations could be excluded as the colony of origin. Although our markers did not prove adequate for assigning individuals within the North Atlantic, this does not mean that a genetic approach may not be valid for other types of markers. Microsatellite markers are found within the nuclear genome and are considered to be neutral (i.e. unaffected by selection), making them ideal for examining population-level processes such as dispersal and changes in the effective number of breeders (see below). However, they may not reflect important adaptive variation, especially if there is mild gene flow or recent selection. A genetic marker that reflects adaptive differences could therefore prove useful for assigning individuals to their population or area of origin. Loci involved in the immune response (such as genes of the major histocompatibility complex, MHC) might be ideal markers, as these loci are highly variable and may be under strong selection given the frequency and intensity of parasitism in colonial seabirds and the potential for local adaptation (McCoy et al.

Table 3. Average pairwise estimates of  $F_{st}$  between common guillemot populations in the North Atlantic (lower matrix) and the p-value associated with each value (upper matrix). Estimates that were significant after Bonferroni correction are given in **bold**. Abbreviations as in Table 1

	HO	RO	GI	LT	SL	SH	SD	HE	GR	NO	FO	SU	F1	WH	SK	HL	CO	LA	CS	SO	CF	WB
HO	0.76569	0.91884	0.02836	0.11017	0.30374	0.20416	0.2463	0.4303	0.28315	0.28045	0.21614	0.34611	0.01977	0.13739	0.25508	0.06064	0.4374	0.13358	0.00879	0.24801	0.03652	
RO	0.0016	0.79777	0.02139	0.19666	0.43797	0.05127	0.61002	0.0331	0.43035	0.56987	0.66357	0.90731	0.37642	0.05603	0.78945	0.0633	0.81983	0.28507	0.19253	0.39026	0.00307	
GI	-0.0059	-0.0027	0.18737	0.35094	0.1679	0.31114	0.38152	0.06879	0.37832	0.26364	0.66061	0.89359	0.7726	0.11194	0.64096	0.15894	0.69497	0.27239	0.14868	0.30124	0.02367	
LT	0.0127	0.011	0.10928	0.29024	0.04426	0.23645	0.00547	0.6319	0.02582	0.0052	0.63002	0.0867	<b>0.0001</b>	0.05765	0.27457	0.39821	0.05586	0.01509	0.02572	0.09838		
SL	0.0214	0.0082	0.0052	0.0114	0.92401	0.24423	0.12506	0.01672	0.3179	0.4497	0.18217	0.27822	0.5693	0.10393	0.29825	0.31624	0.63366	0.4008	0.07365	0.09938		
SH	0.0017	0.0007	-0.0015	0.0022	-0.0076	0.10259	0.33854	0.16043	0.27244	0.59022	0.02238	0.4666	0.47695	0.01243	0.60454	0.39849	0.89867	0.26465	0.03394	0.08294		
SD	0.0127	0.0163	0.0012	0.0128	0.0114	0.0075	0.17366	0.54952	0.51207	0.00295	0.003	0.22208	0.14649	0.03551	0.20127	0.0676	0.0824	0.06753	0.03244	0.0108		
HE	0.0045	0.0011	0.0032	-0.0014	0.0152	-0.0006	0.0145	0.09903	0.65641	0.51304	0.54803	0.80761	0.59751	0.11778	0.72829	0.45943	0.80257	0.3407	0.06112	0.5209		
GR	-0.0041	0.0095	0.0045	0.0146	0.0186	0.0003	0.0022	0.006	0.16605	0.00862	0.00015	0.0499	0.0168	0.00024	0.0066	0.03276	0.13119	0.002	0.00126	0.00083		
NO	0.0044	0.0012	0.0001	-0.0058	0.0018	-0.0004	0.0024	-0.0015	0.0048	0.61711	0.26325	0.57641	0.20714	0.16033	0.89867	0.13425	0.92707	0.31019	0.0318	0.30861		
SU	0.0089	0.0029	-0.0021	0.012	-0.0023	-0.0066	0.0093	0.0029	0.01	0.0003	0.0026	0.002	0.0026	0.62584	0.81695	0.59574	0.31153	0.95469	0.11465	0.56192		
F1	0.0092	-0.0033	-0.0036	0.0077	0.0087	0.0018	0.0169	-0.0017	0.0186	-0.0016	-0.0026	0.0026	0.28322	0.54729	0.01654	0.81469	0.30117	0.95492	0.62287	0.14005		
WH	0.0227	0.0083	0.0033	0.0091	-0.0003	-0.0027	0.0154	0.0043	0.0161	0.0031	-0.0049	0.0006	0.0079	0.63937	0.18128	0.98973	0.32296	0.76708	0.40603	0.39086		
SK	0.0106	0.0117	-0.0014	<b>0.0295</b>	0.0136	0.0114	0.0086	0.0106	0.0173	0.0092	0.0016	0.001	0.0027	0.0191	0.00496	0.26024	0.08919	0.98614	0.78212	0.15029		
HL	0.0062	0.0000	-0.0036	0.0091	0.0067	-0.0001	0.0116	-0.003	0.0133	-0.0025	-0.0039	-0.0055	-0.0045	0.0047	-0.0034	0.66161	0.03023	0.26543	0.00031			
CO	0.01	0.0062	-0.0076	-0.006	0.0047	-0.0023	0.0011	-0.0029	0.0065	-0.0028	0.0014	-0.0024	0.0004	0.0041	0.0121	0.0003	0.58672	0.9476	0.24202			
LA	-0.0019	-0.0018	-0.0076	-0.0011	0.0004	-0.0007	0.0109	-0.0018	0.0073	-0.01	-0.004	-0.0092	0.0029	0.0056	0.0082	0.0075	-0.0044	0.67721	0.04947			
CS	0.0226	0.0053	0.0077	0.0124	0.0043	-0.0013	0.0135	0.0102	0.0156	0.0069	-0.0015	0.0013	0.0148	-0.0011	0.0272	0.0114	0.0011	0.003	0.16443			
SO	0.0231	0.0025	-0.0003	0.0105	-0.0022	0.0005	0.0088	0.012	0.0182	0.0032	-0.007	0.0005	0.0045	-0.0024	0.0172	0.0047	0.0003	0.0036	-0.0049			
CF	0.0009	-0.0023	-0.0023	0.001	0.0106	0.0033	0.0183	-0.0031	0.0195	-0.0031	0.0056	-0.0031	-0.003	0.0104	0.0086	-0.0079	-0.0002	-0.0097	0.0147			
WB	0.0322	0.0422	0.0293	0.0078	0.0442	0.0297	<b>0.0517</b>	0.0297	<b>0.0586</b>	0.0314	0.0373	0.0285	0.0212	0.0378	0.0568	0.0317	0.0211	0.021	0.049			

2003). Moreover, the results of biometry and ring-recovery analyses following the 'Erika' oil spill show that birds came from a wide area (Cadiou et al. 2004), and thus they stress the need to further explore reliable ways to identify the origin of oiled individuals.

This study provides important information about the genetic structure and functioning of common guillemot populations in the North Atlantic. Our results suggest that the populations of North-East Atlantic may be viewed as a single management unit (Friesen 1997, Hedrick 2001). The weak genetic differentiation at this spatial scale shows that from a genetic perspective the colonies constitute a single, large population, and the application of metapopulation theory to the population structure of common guillemots will undoubtedly be highly informative (Esler 2000). This idea of a single large population supports previous results obtained using mitochondrial markers (Moum et al. 1991, Friesen et al. 1996b, Moum & Arnason 2001) and Friesen's (1997) hypothesis that the geographic scale of population genetic differentiation tends to be greater in species that undergo distant winter migrations or feed offshore compared to species that remain near the breeding colony or feed inshore. Overall, our results suggest that, despite the large number of guillemots killed in the 'Erika' oil spill, little genetic variation was lost and no genetically unique component of this species was harmed. This assumes, of course, that our markers are representative of the overall genetic characteristics of the populations; because our markers are neutral, there may be important selective variation that could not be assessed (see above).

The weak genetic structure of guillemot populations may be explained by several mutually-non-exclusive processes. First, the current populations could have been established from a single refugium after the last quaternary glaciation (Friesen et al. 1996b). Considering the mutation rate of microsatellites ( $10^{-2}$  to  $10^{-5}$ , that is, 1 mutation every 100 to 100 000 replications generation<sup>-1</sup> for a given locus; Jarne & Lagoda 1996), insufficient time may have elapsed since establishment of these populations to observe large genetic differences (Moum et al. 1991). Second, repeated bottlenecks followed by recolonization may contribute to the weak population-genetic structure of this species (Moum & Arnason 2001). This hypothesis was proposed to explain the comparatively low mitochondrial variation for this species in addition to the weakly structured populations (Moum & Arnason 2001). However, in the present study, we report levels of genetic diversity for microsatellite markers that are comparable or higher than those for other colonial seabird species (Burg 2000, Tirard et al. 2002, McCoy et al. in press). Finally, a lack of strong population structure

can be the direct result of intercolony dispersal. Dispersal opposes the effect of genetic drift and tends to homogenize populations (Slatkin 1989). Intercolony dispersal has been assumed for some common guillemot populations in which colony size increased too quickly to be explained solely by the intrinsic growth rate (Tuck 1961, Nettleship & Evans 1985, Lloyd et al. 1991). Resightings of ringed birds have also suggested that auks, including common guillemots, can disperse between breeding colonies at large spatial scales (Harris & Wanless 1991, Halley & Harris 1993, Lyngs 1993, Kampp & Falk 1998, Harris & Swann 2002). Immigrants could be attracted by high population densities that may indicate a strong survival rate, high reproductive success and better habitat quality (Nettleship & Evans 1985, Lloyd et al. 1991). Although until the last decade it was assumed that common guillemots were highly philopatric (Tuck 1961, Cramp 1985, Hudson 1985), our results suggest that there may be enough large-scale gene flow to reduce genetic structure. This interpretation corroborates recent monitoring work on the dispersal abilities of this species (Halley & Harris 1993, Lyngs 1993, Harris et al. 1996a, Harris & Swann 2002). Nonetheless, our tests of isolation by distance were significant and thus suggest that dispersal may still be limited at larger spatial scales.

Information on the population functioning of the common guillemot can have important implications for conservation. In particular, relatively high intercolony dispersal would mean that populations may have a high capacity for recovery after local disasters. Population resilience may be a consequence of dispersal in combination with an increased number of first-time breeders due to the availability of vacant breeding sites (Cairns & Elliot 1987, Croxall & Rothery 1991, Wiens 1996). In many seabirds, pre-breeders prospect in their natal colonies, as well as in other colonies, before recruitment (Halley & Harris 1993, Cadiou et al. 1994). Prospecting is now commonly viewed as a behavioural mechanism involved in habitat selection that allows an individual to assess environmental quality in different breeding areas using the presence and success of conspecifics (Boulinier et al. 1996, Danchin et al. 1998, Reed et al. 1999). For species in which individuals are attracted to conspecifics, local extinctions may occur when the size of local breeding groups decreases below a certain threshold (Allee effect). An important question is thus the time it takes for a species to re-colonize an area following local extinction. Depending on the colony of origin of the affected birds, and given the complex processes potentially involved in recolonization, the re-establishment of breeding numbers may be long, despite long-distance dispersal.



Our results underline the need to combine information on the species distributions with a knowledge of the different process involved in the metapopulation dynamics of breeders in order to obtain a comprehensive assessment of the ecological impact of an oil spill on seabird populations. Investigations using model simulations under different scenarios should help to integrate the available information for seabirds sensitive to oil pollution. Such a study has, for instance, been performed to assess the potential impact of long-line fisheries on wandering albatross *Diomedea exulans* populations (Inchausti & Weimerskirch 2002). In the case of oil spills, which often occur when birds are away from the colonies, it is important to consider dispersal and the age structure of the population, along with the degree of subpopulation mixing that may vary with time of year. Such a modelling approach could allow one to use the available information to assess the consequences of a specific oil spill on different seabird populations, but also more generally to determine the scale and type of data needed to assess the potential impact on breeding populations.

The frequency of large oil spills has highlighted the fact that national authorities are often ill-prepared to manage such accidents (Camphuysen & Heubeck 2001, Heubeck et al. 2003). Because spills can have major impacts on the marine environment, as well as on local economies and national politics, it is essential to reinforce preventative international conventions. From a conservation perspective, further work on methods to assess the geographic origins of oiled individuals and on predicting the population-level implications of spills is needed. Information on the population functioning of marine organisms, such as that revealed in the present study, should help us to move in this direction.

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