GEOGRAPHIC ASSIGNMENT OF SEABIRDS TO THEIR ORIGIN: COMBINING MORPHOLOGIC, GENETIC, AND BIOGEOCHEMICAL ANALYSES

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Abstract. Longline fisheries, oil spills, and offshore wind farms are some of the major threats increasing seabird mortality at sea, but the impact of these threats on specific populations has been difficult to determine so far. We tested the use of molecular markers, morphometric measures, and stable isotope (δ15N and δ13C) and trace element concentrations in the first primary feather (grown at the end of the breeding period) to assign the geographic origin of Calonectris shearwaters. Overall, we sampled birds from three taxa: 13 Mediterranean Cory’s Shearwater (Calonectris diomedea diomedea) breeding sites, 10 Atlantic Cory’s Shearwater (Calonectris diomedea borealis) breeding sites, and one Cape Verde Shearwater (C. edwardsii) breeding site. Assignment rates were investigated at three spatial scales: breeding colony, breeding archipelago, and taxa levels. Genetic analyses based on the mitochondrial control region (198 birds from 21 breeding colonies) correctly assigned 100% of birds to the three main taxa but failed in detecting geographic structuring at lower scales. Discriminant analyses based on trace elements composition achieved the best rate of correct assignment to colony (77.5%). Body measurements or stable isotopes mainly succeeded in assigning individuals among taxa (87.9% and 89.9%, respectively) but failed at the colony level (27.1% and 38.0%, respectively). Combining all three approaches (morphometrics, isotopes, and trace elements on 186 birds from 15 breeding colonies) substantially improved correct classifications (86.0%, 90.7%, and 100% among colonies, archipelagos, and taxa, respectively). Validations using two independent data sets and jackknife cross-validation confirmed the robustness of the combined approach in the colony assignment (62.5%, 58.8%, and 69.8% for each validation test, respectively). A preliminary application of the discriminant model based on stable isotope δ15N and δ13C values and trace elements (219 birds from 17 breeding sites) showed that 41 Cory’s Shearwaters caught by western Mediterranean long-liners came mainly from breeding colonies in Menorca (48.8%), Ibiza (14.6%), and Crete (31.7%). Our findings show that combining analyses of trace elements and stable isotopes on feathers can achieve high rates of correct geographic assignment of birds in the marine environment, opening new prospects for the study of seabird mortality at sea.

Key words: assignment methods; biogeochemical markers; bird mortality at sea; Calonectris spp.; longline fisheries; mitochondrial DNA; shearwaters; spatial structure.

INTRODUCTION

Interactions between seabirds and anthropogenic activities frequently result in the death of many individuals. Threats posed by human activities at sea, such as long-lining, offshore wind farms, and oil spills, have increased dramatically over the last decades. Globally, long-lining is one of the world’s most serious threats for pelagic seabirds (Lewison et al. 2005). Seabird interactions with longline fisheries often results in the accidental mortality of many seabirds that are hooked when stealing baits (Brothers et al. 1999). Besides, recent major oil spills have been directly responsible for the deaths of several hundred thousands of birds (Cadiou et al. 2003, Carter 2003, García et al. 2003). In addition, the construction of wind facilities offshore may become Europe’s most extensive technical development in marine habitats, but relevant data on the impact on seabirds are only just starting to become available (Garthe and Hüppop 2004, Hüppop et al. 2006).

Since seabirds are particularly vulnerable to any factor increasing adult mortality, chronic and accidental impacts of human activities at sea can jeopardize the viability of populations (Lebreton and Clobert 1991). Although conservation concerns have mainly focused on the impact of threats at the breeding colonies (Boersma et al. 2002), the exposure of mixing populations to the same threats in specific wintering areas can compromise the viability of multiple populations (González-Solis et al. 2007). Thus, understanding the impact of seabird mortality at sea requires not only quantifying the number of seabirds killed, but also determining their
Assignment methods, whereby individual birds are assigned to a breeding colony, breeding archipelago, or taxon, are only starting to be used. Banded birds can give information about their origin, but recoveries are typically very few, even when some large-scale banding programs have expended enormous efforts (Brown et al. 2004). Nevertheless, intrinsic markers, either biological (e.g., genetic and morphometric variation) or biogeochemical (stable isotope signatures and trace elements concentrations), open new avenues in the assignment of dead birds to their population of origin.

Morphological variation can be used to assess the geographic origin whenever morphotypes are geographically segregated across the breeding range (Webster et al. 2002). Although several studies have shown that the spatial resolution of this approach is often limited (Stratford and Partridge 1996, Cadiou et al. 2003, 2004), examining morphological differences is easy, fast, and inexpensive, and whenever possible should always be considered as a potential tool in assignment studies.

Individual assignment based on genetic data has been extensively used in forensic applications, and their value in the assignment mostly demonstrated (Manel et al. 2005). Most seabird species show strong philopatry and site fidelity (Austin et al. 1994, Bried and Jouventin 2002), which should promote genetic differentiation between populations and thus enhance the potential value of molecular markers for assignment studies. However, recent studies on several seabird species have shown that dispersal plays a major role as a homogenizing force preventing genetic differentiation (see review by Friesen et al. 2007), and thus molecular markers are rendered ineffective (Edwards et al. 2001).

The existence of habitat-specific isotopic signatures also allows the assignment of individuals to their geographic origin (Hobson 2005, Wunder et al. 2005). As tissues incorporate the isotopic content of the prey consumed in a predictable manner, which in turn show spatial structure and pattern in their biogeochemical attributes, these patterns may be used to identify the geographic origin of individuals from any given population. Feather analyses are being extensively used for this purpose, particularly when molting patterns are known, because once formed, feathers become inert, reflecting the elements and the isotopic forms assimilated through the diet when and where they were grown (Chamberlain et al. 1997, Cherel et al. 2000, Becker et al. 2002). Isotopic signatures among different populations can differ for several reasons. Natural isotopic gradients in baseline δ13C have been described in the marine environment (Mehl et al. 2004, Forero et al. 2005, Quillfeldt et al. 2005). In addition, differences in diet or foraging behavior among geographically disjunct seabird populations should result in spatial differences in N and C isotope values of feathers grown under distinct oceanic conditions (Cherel et al. 2006).

A relatively new approach in the assignment of individuals of unknown origin is the use of trace elements. Recent studies suggest their value as intrinsic markers for assignment studies (Szép et al. 2003, Hobson 2005). Whereas isotope composition mainly differs on a continental or a regional scale (>1000 km) (Marra et al. 1998), elemental composition of some animal tissues can vary microgeographically (10–1000 km) among populations at the local scale (Szép et al. 2003). However, whether trace element concentrations vary in accordance with biogeographical gradients is unclear, and data regarding patterns of variation in the marine environment is still lacking.

Overall, despite the potential value of morphology, molecular markers, trace elements, and stable isotopes for the assignment of individuals of unknown origin, several studies showed little or no geographic structure among populations over regional to local scales (Edwards et al. 2001, Wassenaar and Hobson 2001, Cadiou et al. 2004, Riffaut et al. 2005). Therefore, a thorough examination of the potential strengths and weaknesses of the different types of markers is needed. Differences in the accuracy among intrinsic markers and the potential benefit of combining different approaches have been reported (Royle and Rubenstein 2004, Kelly et al. 2005), but comparative studies and further applications are still scarce, particularly in the marine environment.

In the present study we use the Mediterranean Cory’s Shearwater (Calonectris diomedea diomedea; see Plate 1), the Atlantic Cory’s Shearwater (Calonectris diomedea borealis), and the Cape Verde Shearwater (C. edwardsii) as a case study in the analysis of different assignment methods to identify the population of origin of oceanic birds. In this paper we aim first to assess the existence of spatial patterns of variation in the three taxa by using genetic, morphometric, and biogeochemical markers. Second, we evaluate the utility of all four markers as effective tools in assigning Calonectris shearwaters to source populations. Finally, we carry out a preliminary application of the results obtained to assign 41 Cory’s Shearwaters caught by longliners from Catalonian harbors (northeast Spain) operating on the Mediterranean coast.

Materials and Methodology

Study species, study area, and sampling design

Cory’s Shearwaters breed on islands across the Mediterranean from the Iberian coast to the Adriatic and Aegean (mainly C. diomedea diomedea), and in the northeast Atlantic islands, from the Canary to Azores archipelagos (mainly C. diomedea borealis), whereas the Cape Verde Shearwater is an endemic of the Cape Verde Archipelago (C. edwardsii) (Appendix A). Although strong patterns of phylogeographic structure have been described within the genus separating the two Cory’s
subsidiaries and the Cape Verde Shearwater species (Gómez-Díaz et al. 2006), most genetic studies have failed to detect spatial structuring among populations (Randi et al. 1989, Wink et al. 1993, Heidrich et al. 1996, Carneiro da Silva and Granadeiro 1999), but see Rabouam et al. (2000). According to previous genetic studies on the species (Gómez-Díaz et al. 2006), we considered Almeria as an Atlantic Cory’s Shearwater colony within the Mediterranean. The Almeria–Oran Oceanographic Front represents a major oceanographic discontinuity in the Mediterranean, and is the real boundary between Atlantic and Mediterranean surface waters (Beckers et al. 1997). The influence of Atlantic waters entering the Mediterranean would explain the pattern observed for Almeria.

From 2001 to 2005 we collected blood samples, the first (innermost) primary feather, and morphometric measurements from adult birds from up to 21 breeding colonies of the Cory’s Shearwater across the Mediterranean and the northeastern Atlantic, and one breeding colony of the Cape Verde Shearwater (Fig. 1). In addition, we collected tissue samples, the first primary feather and morphometric measurements from 41 Cory’s Shearwaters caught by Mediterranean longliners in 2003 and 2004. Birds were caught during prelaying period (July), incubation (3), chick rearing (3), at the end of the breeding period (October) (26), or the postbreeding period (November) (2). In Cory’s Shearwater the first primary feather is known to be grown at the end of the breeding period, when parents are still feeding their chicks (Monteiro and Furness 1996; J. González-Solís and E. Gómez-Díaz, personal observation).

We considered three spatial scales in the analyses. First, at a local scale, we considered a colony level corresponding to single breeding colonies at a specific island. At a regional scale we considered an archipelago level, grouping several colonies breeding on geographically neighboring islands (i.e., the Balearic, Azores, and Canary archipelagos). Finally, we grouped colonies into the three main taxa involved: the Atlantic and Mediterranean Cory’s Shearwater, and the Cape Verde Shearwater.

**Genetic analyses**

Genetic analyses were performed on 188 Cory’s and 10 Cape Verde Shearwaters from 21 breeding colonies. We chose the mitochondrial control region as a genetic marker to investigate the genetic structure of the species. As no genetic structure was found below the subspecies level (see Results), we also employed ISSR (Inter Simple Sequence Repeats) multilocus fingerprinting to examine separately closer genetic relationships among Atlantic and Mediterranean Cory’s Shearwater breeding colonies.

**DNA isolation, amplification, and sequencing**

DNA was isolated from ethanol-preserved whole blood using the salting-out extraction protocol from Bruford et al. (1998). We amplified a 293-bp fragment of Domain I of the mitochondrial control region of all three Calonectris species using three specific primers that had been designed previously for the species (Gómez-Díaz et al. 2006): either CAL2H (5’CATCCC-ATCCAACCTAAAG3’ or CAL4H (5’AGCCTAT-GTATGGATGTGCA3’). Reaction conditions and automated sequencing were those described by Gómez-Díaz et al. (2006). Representative sequences for each breeding colony are available in GenBank (accession nos. DQ371968-DQ372018).

**Population structure**

We evaluated population genetic structure based on the $F_{ST}$ estimates or Wright’s fixation index of population differentiation using the nested analysis of molecular variance (AMOVA) included in ARLEQUIN (Schneider et al. 2000). The AMOVA treatment reflects the correlation of the haplotypic diversity in a hierarchical analysis (Excoffier et al. 1992). In addition, we performed a neighbor-joining clustering analysis, based on the $F_{ST}$ pairwise genetic distances showing genetic relationships among the Cory’s Shearwater and the Cape Verde Shearwater colonies. To assess the existence of a spatial pattern in the genetic structure of the species, we tested the correlation between genetic distances, measured as $F_{ST}/(1 – F_{ST})$, and geographic distances, calculated as ln(geographic distance by sea) of population pairs (Rousset 1997). We applied Procrustes analysis (least-squares orthogonal mapping) a nonparametric approach for the comparison of the two kinds of nonlinearized data sets, using SPSS 12.0 (SPSS 2003). The method is based on matching corresponding points (landmarks) from each of the two data sets, and provides a measure of fit ($m^2$). The significance test of the $m^2$ statistic was determined by employing a randomization approach to one of the data sets (Protest).

**ISSR fingerprint**

We screened a subset of samples with six RAPD (Random Amplified Polymorphic DNA) and two ISSR (Inter Simple Sequence Repeats) markers. Based on clarity and resolution of fingerprint profiles from this screening, we chose the two ISSR markers ((ACTG (AC)$_7$) and (GTCAAGG (CT)$_6$), which produced scoreable and highly polymorphic PCR products. To avoid intra-assay variability, the amplification protocol was performed below high-stringency conditions and PCR reactions were established according to a standardized protocol (Arribas et al. 1997) with annealing at 65°/50°C (AC/CT primer). We electrophoresed PCR (Polymerase Chain Reaction) products on an 8 mol/L urea/6% polyacrylamide sequencing gel. ISSR markers were detected by silver staining (Bassam et al. 1991). First, we performed duplicate PCR reactions in a control set of individuals to evaluate the reproducibility of ISSR
fragments. We checked repeatability for independent DNA extractions, different PCR reactions, and different PCR runs to explore all possible sources of error, and we excluded for subsequent statistical analyses nonreproducible bands as well as faint bands (Pérez et al. 1998). All the scoring was repeated twice, and to further avoid biased results, analyses were restricted to breeding colonies with a sample size >10 birds (Nybom 2004). For the data analysis, assumptions following analytical procedures by Lynch and Milligan (1994) were applied.

**Biometric analysis**

We performed a clustering analysis on the biometric data to assess the population structure and the degree of morphologic differentiation among Cory’s Shearwater colonies and the Cape Verde Shearwater across its geographical distribution. Birds were sexed using the molecular method described by Fridolfsson and Ellegren (1999). We included six biometric measures: tarsus, wing, maximum head (cranium length plus the bill length) and bill length, bill depth, and bill depth at nostril, of 703 sexed individuals from 19 distinct colonies. To remove the effect of sexual size dimorphism on the assessment of morphometric differences among breeding colonies, morphometric variables of males and females were standardized by subtracting the species mean of each sex. We calculated the Euclidean distance for all pairwise combinations of populations and constructed a cladogram from the similarity matrix using the Neighbor-Joining clustering analysis implemented in the NTSYSpc package version 2.1 (Rohlf 1997). To investigate the existence of spatial gradients in morphology, we tested the relationship between each of the six morphometric measures and the corresponding latitude and longitude coordinates of the sampled colonies by least square and curvilinear regressions using the SPSS 12.0 package (SPSS 2003).

**Isotopic and trace elements analysis**

We analyzed C and N stable isotope signatures and trace element concentration on the first primary feather of 219 individuals from 17 breeding colonies. First, feathers were cleaned from surface contaminants using NaOH and ground to powder using a SPEX 6750 Freezer/Mill (SPEX CertiPrep, Metuchen, New Jersey, USA). To analyze stable isotopes, 0.3 mg of this powder was weighed to the nearest microgram and placed into a Sn (tin) capsule. Samples were oxidized with CuO and CO$_2$/Ag at about 900°C in a Flash EA 1112 Elemental Analyser (CE Elantech, Lakewood, New Jersey, USA) coupled to a pirolizer TC-EA and a breath bench, through an interface Confo III (Finnigan MAT, San Jose, California, USA). NO$_X$ was reduced with Cu at 680°C. The combustion products, N$_2$ and CO$_2$, were dried using MgClO$_4$ and transported to a Delta C Finnigan MAT mass spectrometer (Isotopic...
ratio mass spectrometry, Serveis Científico-Tècnics of University of Barcelona, Barcelona, Spain). International standards were run with each of 12 samples; IAEA CH₃ (87% of C), IAEA CH₄ (42% of C) and USGS 24 (100% of C) for ¹³C and IAEA N1 and IAEA N2 (with 21% of N) and IAEA NO₃ (13.8% of N) for ¹⁵N.

For the analysis of trace elements, 0.1 g of feather powder was digested in 1 mL of nitric acid and 0.5 mL of hydrogen peroxide using teflon bombs for 12 hours at 60°C. The result of the digestion was diluted in 7 mL of distilled water. Analyses were performed using the Atomic Adsorption Spectrophotometry (AAS) technique with a mass Perkin Elmer Optima 6000 Elan Spectrophotometer (Isotopic ratio mass spectrometry, Serveis Científico-Tècnics of University of Barcelona, Barcelona, Spain). Accuracy of analysis was checked by measuring certified reference material (Human Hair CRM 397). To check the reproducibility of the procedure, we included sample replicates as well as negative controls in each set of samples analyzed. In addition, we replicated the analysis in 50 of the samples selected at random. We compared mean breeding colony values of trace element composition between replicates using \( t \) tests for two related samples (SPSS 2003). Only those elements showing significant correlations, \( R^2 > 0.5 \) and no significant differences in mean breeding colony values between the two analyses, were considered for the assignment analyses.

First, to evaluate the degree of structuring among Cory’s and the Cape Verde Shearwater colonies, for both C and N stable isotope and trace elements, we calculated the Euclidean distance for all pairwise comparisons of mean colony values and constructed a cladogram from the similarity matrix using the Neighbor-Joining clustering analysis implemented in the NTSYSpc package version 2.1 (Rohlf 1997). Second, to investigate the existence of spatial gradients, we tested the relationships of carbon and nitrogen isotopic signatures and trace element concentrations with the corresponding latitude and longitude coordinates of the sampled colonies by least-square and curvilinear regression using the SPSS 12.0 package (SPSS 2003).

**Assignments analyses**

Assignment tests on multilocus ISSR data were performed using DOH (Paetkau et al. 1995), a frequency-based program. We split our data set into both baseline data, to calculate population frequency likelihoods, and test data, for the frequency-based assignment, randomly grouping 80% and 20% of the individuals, respectively.

We employed classificatory discriminant analysis (SPSS 2003) to evaluate the ability of morphometrics, stable isotopes, and trace elements in the assignment of birds to their geographical origin. Trace element concentrations were transformed logarithmically to approach normality. We carried out the analysis separately and in combination using the same subset of birds (185 individuals from 15 breeding colonies). For all the analyses, we split the original data set into training data, to build models, and an independent data set, to test the classificatory method, randomly grouping 70% and 30% of the individuals, respectively. In addition, we tested models by jackknife cross-validation. Models were built step by step, including independent variables according to the Wilks’ lambda criterion, and colonies were weighted according to the sample size. Discriminant functions were used to classify the individuals previously excluded in testing for the accuracy of the model. Validation of the model was further examined using samples from three distinct Cory’s Shearwater colonies sampled in previous years and analyzed separately.

After comparing all three assignment approaches (see **Results**), we built a final model using trace elements and stable isotopes on a larger data set, covering almost the entire breeding range of the species, and applied this model to assign 41 bycatch Cory’s Shearwaters.

**Results**

**Genetics**

Clustering analysis on mtDNA data indicated a clear geographical pattern; three groups of colonies corresponded to each of the two geographically isolated Cory’s Shearwater subspecies and the Cape Verde species (Fig. 2A).

AMOVA results suggested strong genetic structuring among the three main taxa (\( F_{CT} = 0.61, P < 0.001 \)). The largest variance component was due to differences among subspecies (60.5%); whereas within each subspecies, among breeding colonies, the variance component was lower (37.2%). However, a second AMOVA, at a local scale, failed to reveal genetic structuring among breeding colonies within either the Atlantic or the Mediterranean subspecies region (\( F_{ST} = 0.040, P = 0.350; F_{ST} = 0.00, P = 0.450 \), respectively).

Procrustes analyses revealed a significant correlation between genetic and geographic distances among colonies considering both the Atlantic and the Mediterranean Cory’s Shearwater subspecies regions (\( m^{12} = 0.63, P < 0.001 \)). However, at a local scale among colonies within each subspecies region, the correlation was lower. Whereas spatial correlation was still significant within the Mediterranean (\( m^{12} = 0.63, P = 0.020 \), significance did not hold within the Atlantic (\( m^{12} = 0.93, P = 0.716 \)).

**Biometrics**

The cladogram represents the similarity pattern of the 19 breeding colonies included in the analysis, considering six biometric measures (Fig. 2B). Three groups were clearly defined, corresponding to each of the three taxa. A first group included populations corresponding to the largest in size, the Atlantic Cory’s Shearwater. All Mediterranean breeding colonies grouped separately corresponding to the intermediate in body size, the
Mediterranean Cory’s Shearwater. The smallest in size, the Cape Verde Shearwater breeding colony, was grouped separately.

Linear regression analyses revealed a significant positive longitudinal pattern within the Mediterranean subspecies in three of six biometric measures for males and females (tarsus: $R^2 = 0.16$, $n = 108$, $P < 0.001$ and $R^2 = 0.10$, $n = 94$, $P = 0.002$; bill depth at nostril: $R^2 = 0.24$, $n = 108$, $P < 0.001$ and $R^2 = 0.13$, $n = 96$, $P < 0.001$; bill depth: $R^2 = 0.13$, $n = 108$, $P < 0.001$ and $R^2 = 0.14$, $n = 96$, $P < 0.001$, for males and females, respectively). For the Atlantic subspecies no significant association was found between any of the six measures and either longitude or latitude (all measures: $R^2$, $0.15$, $n = 224$ and $R^2 < 0.05$, $n = 228$, for males and females, respectively). Both linear and quadratic regression coefficients and $F$ statistics are shown in Appendix D. In a few cases quadratic models fit substantially better than linear models.

Isotopes and trace elements

The cladogram based on carbon and nitrogen signatures grouped most breeding colonies according to their geographical distributions. All Mediterranean breeding colonies grouped together except Tremiti, ungrouped, and Almeria, grouped within the Atlantic cluster. In the Atlantic, northeast and southeast Atlantic breeding colonies grouped separately (Fig. 2C). The similarity tree based on trace elements composition showed no geographical structure among breeding colonies, nor among the three main taxa. Nevertheless, breeding colonies appeared well differentiated and large distances separated each node (Fig. 2D).

Regression models based on stable isotope composition of primary feathers showed significant spatial gradients for both $\delta^{15}N$ and $\delta^{13}C$ values across the breeding range of the study species (Fig. 3; Appendix C). Regression coefficients and $F$ statistics are shown in Appendix D. Least squares regression models, although significant, did not explain much variability, whereas curvilinear models, in particular cubic regressions, better fit the spatial gradients observed (Appendix D). Carbon isotope ratios were associated with latitude ($R^2 = 0.17$ and 0.24; least squares and cubic regression, respectively) (Fig. 3A), decreasing from south (Gran Canaria Island, $\delta^{13}C = -13.34 \pm 0.69$) to north (Menorca Island,
$\delta^{13}C = -16.23 \pm 0.32$), except for the most southern Cape Verde breeding colony, which showed unexpected low values ($\delta^{13}C = -15.66 \pm 0.76$). No significant association was found between carbon isotopic signatures and longitude ($R^2 = 0.06$). At a local scale, within each Cory’s subspecies region, the latitudinal decrease in carbon values was further apparent within the Atlantic ($R^2 = 0.49$ and 0.61; least squares and cubic regression) (Fig. 4A), but also within the Mediterranean ($R^2 = 0.35$ and 0.39; linear and cubic regression).

Globally, nitrogen isotope values in feathers tended to show an inverse relationship with longitude, decreasing from west to east ($R^2 = 0.45$ and 0.56; least squares and cubic regression) (Fig. 3B). Indeed, significant differences in mean nitrogen isotopic signatures split the eastern $C. diomedea diomedea$ ($\delta^{15}N = 10.57 \pm 1.34$) from the western $C. diomedea borealis$ breeding colonies ($\delta^{15}N = 12.97 \pm 0.74$; $t = -15.36$, $P < 0.001$). However, the colony from Tremiti showed greater nitrogen isotope values ($\delta^{15}N = 12.28 \pm 1.03$) than those reported within the entire Mediterranean ($\delta^{15}N = 10.57 \pm 1.34$). No significant association was found between nitrogen isotope values and latitude ($R^2 = 0.07$). Considering each subspecies range separately, nitrogen isotope values tended to decrease with longitude within the Mediterranean ($R^2 = 0.23$ and 0.64; least squares and cubic regression) (Fig. 4B), whereas no pattern was apparent within the Atlantic ($R^2 = 0.07$ and 0.08; linear and cubic regression; Appendix D).

Results from ANOVA showed significant differences in trace element composition (Na, Mg, Mn, Ba, S, Zn, Al, P, Ca, Cr, Rb, Sr, Sb, La, Ce, Pr, Nd, Hg, and U) of primary feathers across the study species range (all $P < 0.001$) (Appendix B). Most elements showed weak associations with either latitude or longitude (least squares and cubic regression coefficients, $R^2$, ranging from 0 to 0.20). Only a few elements showed $R^2 > 0.20$ (Appendix D). That is, Mg, Sr, and U concentrations varied with longitude across the breeding range ($Mg, R^2 = 0.20$, Sr, $R^2 = 0.23$, and U, $R^2 = 0.34$; for cubic regression), and Hg was mainly associated with latitude ($R^2 = 0.28$ and 0.37; least squares and cubic regression). No significant spatial correlations were found in Hg levels within the Atlantic or the Mediterranean. In contrast, Mg, Sr, and U were notably associated with longitude within the Mediterranean subspecies range (Mg, $R^2 = 0.27$ and 0.42, Sr, $R^2 = 0.25$ and 0.45, and U, $R^2 = 0.35$ and 0.44, for least squares and cubic regression, respectively.) (Regression coefficients and $F$ statistics are shown in Appendix D.)

**Geographic assignment**

Genetic assignment based on ISSR data failed to assign individuals to breeding colonies. In the randomization procedure only 62 out of 313 Cory’s Shearwaters were assigned to the putative colony. The rate of correct assignment was 22.4% among breeding colonies for the Mediterranean Cory’s Shearwater subspecies, whereas in the Atlantic only 16.8% of individuals were correctly classified. For the independent data set 11.5% and 11.9% of individuals were correctly assigned among the Atlantic and the Mediterranean subspecies, respectively.

The discriminant analysis based on trace elements composition achieved the best rate of correct assignment at both colony and archipelago levels, whereas discriminant models based only on either
morphometric measures or stable isotopes just showed

high rates of correct assignment among taxa (Table 1). In both independent data set and independent analysis, the ability to correctly classify individuals dropped noticeably in all three assignment approaches when used separately; the more detailed the scale of the analysis, the larger the drop (Table 1). Combining different approaches substantially improved the rate of correct assignments at both colony and archipelago levels in most cases. Models based on isotopes and trace elements, as well as models including those with morphology, achieved the best assignment results. Cross-validations, independent data, and independent analysis further confirmed the ability of the combined models, based only on isotopes and trace elements or together with morphology, to discriminate among geographic origins (Table 1).

A final model built on 219 individuals from 17 breeding colonies combined C and N isotopic signatures and trace element concentrations, to assign bycatch birds whose origin was unknown. Discriminant analysis achieved rates of correct assignment >80% for all the

Table 1. Classification rates obtained for each assignment method at each scale of analysis when used alone and in conjunction.

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Notes: Correct assignments (percentages) are shown for training data, test data, jackknife cross-validation, and independent analysis. Independent test data were the 30% of the data randomly excluded prior to building the discriminant model. Independent test analysis corresponded to samples collected in previous years and analyzed separately.
Mitochondrial DNA analyses revealed a strong genetic structuring among the three main taxa, agreeing with their geographically segregated distributions (Fig. 2A). However, the Mediterranean colony of Almeria was placed within the Atlantic cluster. This result agrees with previous genetic studies on the species, pointing out the Atlantic subspecies identity of Almeria (Gómez-Díaz et al. 2006).

The mitochondrial control region is a rapidly evolving locus extensively used for population-level studies (see Pearce 2006), and its potential utility to discriminate among geographic origins has been previously shown in a number of species (Lovette et al. 2004). However, this gene appeared ineffective for the geographic assessment of Cory’s Shearwaters beyond the taxon level. Similarly, although ISSR multilocus fingerprints revealed substantial genetic variability, assignment tests failed to classify individuals to the breeding colony. The low level of genetic structuring observed for both molecular markers explains the weak spatial resolution and the low rates of correct assignments. Lack of genetic structure has been repeatedly reported for the species using different genetic markers (Randi et al. 1989, Wink et al. 1993, Heidrich et al. 1996, Carneiro da Silva and Granadeiro 1999, Gómez-Díaz et al. 2006). Analyses based on microsatellite fingerprints found genetic structuring at lower levels, both among and within breeding sites, but failed to detect a correlation between spatial and genetic patterns of variation (Rabouam et al. 2000).

**Discussion**

**Genetics**

Mitochondrial DNA analyses revealed a strong genetic structuring among the three main taxa, agreeing with their geographically segregated distributions (Fig. 2A). However, the Mediterranean colony of Almeria was placed within the Atlantic cluster. This result agrees with previous genetic studies on the species, pointing out the Atlantic subspecies identity of Almeria (Gómez-Díaz et al. 2006).
stable isotopes signatures separated most breeding colonies according to their geographic distributions, although to a lesser extent than the biometric and genetic structuring. The influence of Atlantic waters entering the Mediterranean would explain the isotopic similarity of the three most western Mediterranean colonies (i.e., Murcia, Almeria, and Chafarinas) to those of the Atlantic (Fig. 2C). In contrast, although showing the largest distances compared to any other approach, no geographical structure among breeding colonies was evident for trace elements.

In the present study, both $\delta^{15}$N and $\delta^{13}$C values of the first primary feather varied across the breeding range of Cory’s Shearwaters and differed from those for the Cape Verde Shearwater. Since the first primary feather is grown at the end of the breeding period, we can expect the isotopic signatures of this feather to reflect those from the prey consumed around the breeding area. However, evidence of biogeographic patterns of stable isotope values in marine ecosystems is still scarce, and several studies showed large differences in the degree of geographic variation of isotope biomarkers, depending on the spatial scale of analysis (Chamberlain et al. 1997, Marra et al. 1998, Wassenaar and Hobson 2001, Wunder et al. 2005). Whereas most studies successfully applied C and N isotope ratios to discern geographically distinct subspecies, most failed to discriminate among geographic origins at regional and local scales (see Hobson 1999 for a complete review). In line with results found in other studies in both terrestrial and marine ecosystems, we found feather isotope values of C and N to vary by latitude and longitude, respectively. However, the strength of these patterns depended on the geographic scale considered (see Figs. 3, 4). The geographic break in isotope values between Atlantic and Mediterranean regions effectively separated the two subspecies. However, isotope geographical gradients within each region, although eventually significant, were too weak to reliably assign the origin of birds at local scales. This inaccuracy probably resulted from variance associated with isotope values at a single location, or from similar isotopic baselines values at different areas (Royle and Rubenstein 2004). Alternatively, similarities in isotopic values among neighboring colonies may simply result from birds sharing the same foraging areas. During the chick-rearing period, when the first primary feather is grown, shearwaters use a flexible foraging strategy, i.e., they forage close to the colonies, mainly to feed the chicks, but far offshore for self feeding (Granadeiro et al. 1998, Cherel et al. 2005), potentially overlapping with shearwaters breeding in other islands.

However, sharing foraging areas of neighboring islands cannot solely explain the lack of spatial resolution of isotopic values, because this should then hold true for the genetic analyses, the Mediterranean colony of Almeria intermingled with the Atlantic cluster. Concerning the spatial pattern of variation in morphology, we found a longitudinal geographic gradient in body measurements within the Mediterranean subspecies as a slight increase in body size from east to west Mediterranean subspecies, a biometric pattern previously recognized by other authors (Massa and Lo Valvo 1986, Granadeiro 1993). This pattern is probably related to the complex longitudinal oceanographic subzonation within the Mediterranean basin. Specific marine habitat characteristics of different areas within the Mediterranean, such as differences in sea surface temperature, wind speed, and primary production, probably promoted divergence in body size through local adaptation of populations (Zotier et al. 1999). On the contrary, no spatial pattern was apparent within the Atlantic, although some authors have recently reported slight latitudinal differences from the Canary to Azores Islands (Thibault et al. 1997, Gómez-Díaz et al. 2006).

Morphological measures could be used to reliably assign most birds to the taxon (species or subspecies), provided that the sex of the birds was previously determined, since Cory’s Shearwaters show an appreciable sexual size dimorphism. However, assignment tests based on morphological data failed to identify the geographic origin of Cory’s Shearwaters at their breeding colony. Recent studies on the seabird genus Puffinus sp. showed that only species with widespread distributions would exhibit noticeable geographic variation in their morphometrics (Bull et al. 2005). Consequently, data supplementary to the more traditional biometric approach, such as isotopes, trace elements, or genetic markers, are probably necessary for the geographic assignment of most seabird species.

### Table 2. Extended.

<table>
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<th>Assigned (%)</th>
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trace element composition of these feathers. Instead, we found specific trace element profiles associated with each breeding location, but in contrast with stable isotopes, we could not detect latitudinal or longitudinal patterns in feather composition (Fig. 2D). Differences in elemental composition of feathers allowed us to discriminate the origin of birds at different geographic scales, that is, from specific breeding colonies (77.5% of correct assignments), considering archipelagos (75.2%), and identifying taxa (89.9%). However, correct classification at the colony and archipelago levels decreased to 62.5% and 69.6%, respectively, when we classified 20% of the samples randomly excluded from the analyses prior to building the discriminant model. The model performed even worse for classifying an independent set of samples collected in previous years and analyzed separately, dropping to 23.5% and 41.2% correct classification. These results illustrate the limitations of these models to classify samples other than those used to build them. The drop in the classification rates of samples collected in previous years may reflect interannual variability in trace element composition. In addition, the lack of spatial gradients in trace elements composition across the breeding range of the species underlies the need for a geographically widespread sampling design. Sampling discontinuities and missing populations would lead to inaccuracies in the assignment of individuals to specific locations. In addition, misclassifications in the validation methods among, i.e., Cape Verde and Canary/Azores missassigned birds (Table 2), would be explained because baseline levels of the three areas are similar. In this case, population trees for any intrinsic marker (Fig. 2) can help us to infer whether the most likely reason for the missassignments is a close similarity among colonies. Still, trace element composition was the intrinsic marker showing the best assignment rates, suggesting that such composition strongly depends upon the microgeographical location of the breeding site. This probably reflects the importance of local conditions and the similarity in foraging areas of birds breeding in the same colony (Szép et al. 2003), although local atmospheric deposition on feathers might also explain these differences to some extent (Breitburg and Riedel 2005).

The combination of several methods enhanced the reliability of the classificatory method both across the different spatial scales (86.0%, 90.7%, and 100.0%, from colony to taxon level), and for all validation methods (e.g., at the archipelago level, 82.1%, 94.1%, and 79.8% for independent data, independent analysis, and jackknife cross-validation, respectively). The benefits of using multiple intrinsic markers has been demonstrated previously (Royle and Rubenstein 2004), and our results highlight that a combined approach should be used when possible.

**Bycatch birds**

The preliminary application of the discriminatory model on Cory’s Shearwaters caught by Mediterranean
longliners identified three source breeding sites in the Mediterranean (Menorca, Crete, and Ibiza islands) and one from the Atlantic (Graciosa Island). Ancillary information on the capture date, location, and numbers of dead birds recovered may help to contrast model predictions and to test ecological hypotheses. Interestingly, both Menorca and Ibiza breeding colonies are within the operating area of Catalonian longliners and represent major breeding colonies for the species in the central Mediterranean (Thibault et al. 1997). Furthermore, most of the birds assigned to Crete were caught out of the breeding period (86.0%), either during the prelaying period (i.e., April and May) or during the postnuptial migration (i.e., October and November), when birds have to cross the operating area of the Catalonian longliners returning from or going to the wintering grounds in the subSaharan coast (Ristow et al. 2000). Finally, although we could expect all individuals to belong to the Mediterranean subspecies, we cannot completely rule out the Atlantic identity of some specimens, since there is evidence of Atlantic Cory’s Shearwaters entering the Mediterranean (Lo Valvo and Massa 1988, Sánchez 1997, Thibault and Bretagnolle 1998, Martínez-Abraín et al. 2002). However, those birds were caught in May and August, within the breeding period of the species, and their biometric measurements do not support their Atlantic identity. As mentioned above, ghost populations resulting from incomplete sampling designs would force the model to assign birds elsewhere. In the present study, although the sampling design covered almost the whole breeding range of the species, a few central Mediterranean populations were missed (mainly Zembra, Malta, Corsica, and Sardinia), which may partly explain assignment errors when applying the model.

Conclusions

Overall, the effectiveness of the four types of biomarkers evaluated, i.e., genetic, morphometric, trace element, and isotopic traces for the geographic assignment of Cory’s Shearwaters, varied with the spatial scale of analysis. The application of intrinsic markers depends on there being enough differences among potential source breeding colonies to unequivocally assign individuals to their source. Which method should be applied also depends on the level of spatial resolution desired and the ancillary information we have, such as the sex of the bird (Fig. 5). Whereas the identification of Atlantic and Mediterranean Cory’s Shearwater subspecies and the Cape Verde species was successfully achieved for all four types of markers, large differences arose in the correct assignment of the breeding colonies among these methods. Both morphometric and genetic approaches appeared ineffective at discriminating among geographically close breeding sites. With respect to biogeochemical markers, stable isotopes showed some latitudinal and longitudinal trends, but the predictive power of these trends to classify individuals among putative sources was rather weak. In contrast, trace elements appeared more sensitive to microgeographical differences than were stable isotopes, and allowed the best rate of correct assignment of feathers at local scales. However, trace elements did not show geographical gradients, and thus ghost populations, the result of incomplete
sampling designs, can force the model to erroneously assign birds to very distant colonies rather than identifying their true origin. Adding isotopes to models based on trace elements did not enhance the rate of successful assignment of the training data, but substantially improved classification rates of the independent data sets, increasing the robustness and accuracy at all the spatial scales of analysis. Determining the origin of dead individuals is important for any attempt to assess the potential impact of human activities, such as oil spills, long-line fishing, or offshore wind farms, responsible for an increase in seabird mortality at sea. Our study shows that it is feasible to use biogeochemical markers of seabird feathers, such as trace elements and stable isotopes, to identify the impact of such activities on specific breeding seabird populations at a local scale.

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APPENDIX A

Geographic location of Cory’s and Cape Verde Shearwaters and sample sizes for each analysis (Ecological Archives A017-058-A1).

APPENDIX B

Morphological measurements for all Calonectris breeding colonies included in the biometric analysis (Ecological Archives A017-058-A2).

APPENDIX C

Mean and standard deviation of stable isotopes signatures and trace elements concentrations of Cory’s Shearwater and the Cape Verde Shearwater breeding colonies (Ecological Archives A017-058-A3).

APPENDIX D

Regression models on the latitudinal and longitudinal gradients in stable isotopes signatures, trace elements concentrations, and morphometric measurements (Ecological Archives A017-058-A4).