Inter-colony movement of pre-breeding seabirds over oceanic scales: implications of cryptic age-classes for conservation and metapopulation dynamics

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(A) RUNNING TITLE

Inter-colony movement of pre-breeding seabirds

(A) ABSTRACT

(B) Aim

Demographic linkage between subpopulations plays a critical role in population processes. Metapopulation dynamics, however, remains one of the most poorly understood aspects of population biology. This is especially true for small, pelagic seabirds because their discrete subpopulations are located on offshore islands, separated by vast areas of open-ocean making monitoring logistically challenging. Seabird populations often contain large numbers of immature pre-breeders that may be important for subpopulation connectivity and demography, but are poorly studied. Here we provide evidence for inter-colony movement of pre-breeding Leach's storm-petrels *Oceanodroma leucorhoa* among three colonies spanning the North Atlantic Ocean. We discuss their influence on metapopulation dynamics and the extinction risk of a subpopulation under threat from extreme predation.

(B) Location

North Atlantic Ocean Islands (Scotland, Canada & Iceland)

(B) Methods

We use a novel application of Bayesian stable isotope mixing models to infer recent movement of pre-breeding birds between three major breeding populations in the North Atlantic. Carbon and nitrogen stable isotope values from breeding birds (central place foragers) sampled at each colony were used as model sources and pre-breeding birds as model mixtures.

(B) Results

Of 134 pre-breeding Leach's storm-petrels sampled at three colonies across the North Atlantic, 5 had isotope mixing model estimates dissimilar (<25%) to their colony of capture and were instead isotopically similar to another breeding region. Nineteen further prebreeders had highly mixed signatures (<50% for the colony of capture) indicating possible recent movement among colonies.

(B) Main Conclusions

Our findings provide evidence for inter-population connectivity of pre-breeding Leach's storm-petrels among colonies spanning the North Atlantic. These results highlight the significance of cryptic young age-classes in metapopulation dynamics and the demographic processes. Moreover, they provide us with a better understanding of how one subpopulation remains extant, despite experiencing extreme predation rates.

(B) Keywords

Extinction risk, dispersal, Leach's storm petrel, metapopulation, predation, stable isotopes.

(A) INTRODUCTION

Highly fragmented populations are reliant upon dispersal to maintain demographic and genetic connectivity (Bowler & Benton, 2005; Lowe & Allendorf, 2010), yet isolation of these subpopulations (demes) can make them vulnerable to extinction (Matthiopoulos et al., 2005). When sufficient dispersal occurs between subpopulations to influence local demographics, but subpopulation independence is maintained, this can be considered a metapopulation (Hanski, 1998). Estimating inter-population connectivity is essential for predicting population and metapopulation dynamics, as well as the potential for subpopulations to buffer the effect of environmental change (Kokko & López-Sepulcre, 2006), but remains one of the most poorly understood processes in population biology. For species with delayed maturation, there is a paucity of empirical data on the movement of prebreeding individuals, although prospecting behaviour is known to play an important role in breeding location decision making (Dittmann et al., 2005; Caut et al., 2008; Robinson & Beckerman, 2013). This knowledge gap is problematic because for long-lived iteroparous species immatures may represent a significant proportion of the fully grown population (Lebreton et al., 2003), play an important role in the demographic process (Votier et al., 2008) and also have a higher dispersal propensity than breeders (Lebreton et al., 2003; Votier et al., 2008).

Many colonial nesting seabirds breed under conditions conducive to the formation of metapopulations. Breeding colonies are discrete subpopulations, separated by a matrix of unsuitable habitat (i.e. the open ocean), and seabirds have excellent dispersal capabilities to aid connectivity (Inchausti & Weimerskirch, 2002; Breton *et al.*, 2006). However, the extent to which seabird colonies function as metapopulations is poorly understood (Oro, 2003). This arises in part because of the logistic constraints of studying (often) remote colonies, comprising large numbers of individuals, separated by vast distances at sea. Moreover, because fidelity to breeding locations is high among most seabirds (Schreiber & Burger, 2002), pre-breeding rather than breeding birds are likely to contribute most to inter-colony connectivity. Seabird populations can consist of large numbers of young pre-breeders (>50% of the population (Klomp & Furness, 1992)), but due to the fact that they rarely come to land (where the majority of seabird research occurs) they lead cryptic lives (Votier *et al.*, 2011). Previous research using capture-mark-recapture techniques suggests pre-breeding seabirds play an important role in metapopulation dynamics (Inchausti & Weimerskirch, 2002;

Lebreton *et al.*, 2003; DeMots *et al.*, 2010; Turner *et al.*, 2010), but this process is virtually unknown for species where marking at birth and subsequent re-sighting is logistically unviable. Stable isotopes signatures of avian tissues are frequently used to investigate migratory connectivity and origins of terrestrial birds (Wunder *et al.*, 2005), and have also been combined with other biochemical markers to assign seabirds to colony of origin (Gómez-Díaz & González-Solís, 2007). However, these rely upon (and are limited by) a precise knowledge of the timing and pattern of moult in each species, and if the main moult occurs outside the breeding season (which is the case for many taxa) then analysis of feather isotopes is not appropriate for investigating the movements of pre-breeders. Here we propose an application of analysing stable isotope signatures in blood to test for inter-colony movement of pre-breeding seabirds at an oceanic scale.

Our study focusses on a species of conservation concern in the EU, the Leach's storm-petrel Oceanodroma leucorhoa (Vieillot). This is a small (<45g), burrow-nesting Procellariiforme that breeds at a small number of widely distributed colonies, separated by large tracts of open ocean, across the North Atlantic and Pacific. At the second largest colony in Europe on St Kilda, Scotland, a ~54% decline in breeding Leach's storm-petrels from 90,000 individuals in 1999 to 41,000 individuals in 2006 (Newson et al., 2008), raised concerns for the viability of this regionally important population. This reduction is partly attributable to predation by great skuas Stercorarius skua (Brunnich), yet the estimated consumption of ~21,000 Leach's storm-petrels year⁻¹ by skuas exceeds the observed decline (Phillips et al., 1999; Miles, 2010). One explanation for this disparity is that prospecting pre-breeders or breeding immigrants from other very large Leach's storm-petrel colonies in the North Atlantic may be buffering the impacts of skua predation (Votier et al., 2006; Bicknell et al., 2012). Over 5 million pairs of Leach's storm petrel breed in Iceland and eastern Canada (Bicknell et al., 2012), indicating huge numbers of potential prospectors in the North Atlantic. Fidelity to breeding locations is high (Blackmer et al., 2004), but recent work has revealed genetic homogeneity among North Atlantic colonies suggesting the occurrence of effective natal dispersal (Bicknell et al., 2012).

We used stable isotopes in blood to test for inter-population connectivity of pre-breeding Leach's storm-petrels spanning >3000km in the north Atlantic. This approach provides answers to questions on an immediate ecological time scale inaccessible using other techniques (e.g. genetic markers, but see; McCoy *et al.*, 2005), but is critical to assess current metapopulation dynamics and conservation threats. First we measured stable isotopes in red blood cells (RBC) of breeders from three colonies representing the main North Atlantic population centres, to check for isotopic differences among regions, in tandem with confirmatory data on zooplankton from the same regions. Second, we used a stable isotope mixing model (SIAR) to estimate the probability of pre-breeders moving among colonies and therefore allowing an assessment of the importance of this age-class in maintaining colony connectivity at ocean-basin scales.

(A) METHODS

Leach's storm-petrel samples

Leach's storm-petrels were sampled from Gull Island, Canada $(47^{\circ}15^{\circ}N, 52^{\circ}46^{\circ}W)$ and St Kilda, Scotland $(57^{\circ}49^{\prime}N, 08^{\circ}35^{\prime}W)$ in August 2008, and Vestmannaeyjar, Iceland $(63^{\circ}25^{\prime}N, 20^{\circ}17^{\prime}W)$ in August 2009 (Fig. 1a). Between 10-20µl of blood was collected from the brachial vein of breeders in burrows (incubated an egg or brooding a chick), as well as prebreeders caught in mist-nets at night under appropriate regional licences. To ensure the status of pre-breeders only birds that met all of the following criteria were included: (1) caught away from the main breeding areas; (2) attracted by a chatter call lure played on speakers close to the mist-net; (3) no regurgitation of prey items when caught or handled; and (4) had no obvious brood patch. Blood was separated into plasma and RBC using a centrifuge within 2-3 hours of sampling, and then freeze-stored until analysis.

(B) Stable isotope analysis

Carbon and nitrogen stable isotopes show naturally occurring gradients that can be used to infer movement at the individual-level (Hobson, 2007). Carbon (δ^{13} C) values are representative of the primary carbon source and show a number of spatial gradients in marine ecosystems (such as inshore vs. offshore, pelagic vs. benthic, and latitude (Hobson, 2007)). Nitrogen ($\delta^{15}N$) values show a consistent step-wise enrichment by trophic level and also reflect the nitrogen pool supporting primary producers at the base of different food webs (Owens, 1987; Jennings & Warr, 2003). We used $\delta^{15}N$ and $\delta^{13}C$ values of RBC because this tissue typically reflects diet ~3-4 weeks prior to sampling (Bearhop et al., 2002) and because there is little confounding influence of lipids (Cherel et al., 2005). Breeding adult Leach's storm-petrels are constrained to tend their eggs or chicks (return to the nest every 1-4 days; Ricklefs et al., 1985) with an estimated foraging range over 200km (Hedd et al., 2009), such that δ^{13} C and δ^{15} N values represent the region close to the colony of capture. Pre-breeders are not constrained in the same way and therefore, if they move between different colonies, may have mixed isotope signatures. We used the Bayesian stable isotope mixing model SIAR (Parnell et al., 2010) to calculate the median probability of pre-breeder isotope signatures attributable to the three colonies of capture, where breeder isotopes are the sources and prebreeders the mixtures. The sources and mixtures were therefore derived from the same trophic level obviating the need for trophic enrichment factors in the model. We assume that any potential differences in assimilation efficiency or physiology between breeders and immature birds did not influence the relationship between isotope values in prey and blood.

(B) SIAR mixing model parameters and performance

To provide an independent test of regional isotopic distinction (not used in any model), we analysed zooplankton samples collected by the Continuous Plankton Recorder Survey (CPR - managed by the Sir Alister Hardy Foundation for Ocean Science) in oceanic waters representative of the three regions. This was not to directly reflect the diet of Leach's storm-petrels (although they do consume some zooplankton (Hedd & Montevecchi, 2006)) but instead to determine whether regional differences in the stable isotope values of breeder RBC are consistent at lower trophic levels within the same year. Adult life stages (CV and CVI) of *Calanus helgolandicus* and *C. finmarchicus* were removed from 27 formalin preserved "silks" (collection mesh representing 10km towed transects) sampled on 5 survey routes (2 in Canadian region, 2 in Icelandic region and 1 in Scottish region), which crossed oceanic regions representative of each colony during July 2008 (Fig. 1a; Table S1). Isotopic values from multiple CPR routes sampled within the Canadian and Icelandic regions were similar and therefore combined into a value for each region (Fig. S1).

Key to mixing model efficacy is accurate characterisation of potential sources (prey samples in diet reconstruction, but colony-specific regions in this instance). Independent data from CPR samples and the literature were used to check our source coverage assumptions. It is also desirable for the mixture values (in this case pre-breeders) in mixing models to lie within the fuzzy convex hull of the source isotope values (breeders), and this was checked using an iso-space plot (Parnell *et al.*, 2012).

To assess SIAR model performance and sensitivity to a small sample size of breeders at one colony (Gull Island, Canada), we re-ran the model using estimates for Gull Island based on subsampled data. We replicated the small sample size for each of the other two sources by randomly sampling 4 δ^{13} C breeder values from their ranges, repeated 10 times to produce a range of possible means for each source (colony). The difference between the lowest mean of the range and the original source mean (all the samples) was averaged from the two sources and then subtracted from the original Gull Island mean. This produced an estimated mean to be use in a sensitivity analysis (see Supplementary data, Dryad repository; DOI: doi:10.5061/dryad.q2610). The average of the St Kilda and Vestmannaeyjar δ^{13} C standard deviations was used as the estimated standard deviation. This process was repeated for the δ^{15} N data. These estimates represent a conservative scenario in terms of source differentiation and variance (i.e. by bringing the sources closer together and increasing the variance; see Fig. S1).

(B) Stable isotope sample preparation

All samples were freeze-dried, homogenised and ~0.7 mg weighed into a tin cup for analysis. Each zooplankton sample (tin cup) consisted of approximately 10-20 individual animals. Metabolic synthesis of lipid discriminates against ¹³C (DeNiro & Epstein, 1977) and generally yields low δ^{13} C values in tissues with high lipid content when compared with an organisms' dietary input (Tieszen *et al.*, 1983). Therefore, because of highly variable lipid content between *C. helgolangicus* and *C. finmarchicus* (Ventura, 2006), lipids were extracted prior to analysis. Inorganic carbonates were also removed from these samples, as these tend to increase δ^{13} C and do not reflect the organisms dietary intake (DeNiro & Epstein, 1978). Extraction and removal methods, as well as correction for formalin preservation, are described in detail elsewhere (Bicknell *et al.*, 2011).

Analyses were conducted at the East Kilbride Node of the Natural Environment Research Council Life Sciences Mass Spectrometry Facility via continuous flow isotope ratio mass spectrometry using a Costech (Milan, Italy) ECS 4010 elemental analyser interfaced with a Thermo Electron (Bremen, Germany) Delta XP mass spectrometer. Isotope ratios are reported as δ -values and expressed as ∞ according to the equation:

 $\delta X = [(R_{sample}/R_{standard})-1] \times 1000$ (Eq 1) where; X is ¹³C or ¹⁵N and R is the corresponding ratio ¹³C/¹²C or ¹⁵N/¹⁴N and R_{standard} is the ratio of the international references PDB for carbon and AIR for nitrogen. The standard deviation of multiple analyses of an internal gelatine standard in each experiment was better than 0.2‰ for δ^{15} N and δ^{13} C. All statistics and models were run in R version 2.15.1 (R Development Core Team, 2011).

(A) **RESULTS**

(B) Isotopic differences among regions (sources)

Breeding Leach's storm-petrels were isotopically segregated among colonies (MANOVA, Pillai, $F_{2,31} = 50.86$, P <0.0001). Univariate analysis revealed δ^{15} N was significantly different between all colonies (ANOVA, $F_{2,31} = 55.18$, P <0.0001) and δ^{13} C was significantly different between Vestmannaeyjar, Iceland and the other colonies (ANOVA, $F_{2,31} = 46.05$, P <0.0001; St Kilda, Scotland, t = 9.26, P < 0.0001; Gull Island, Canada, t = 5.91, P <0.0001), while Gull Island and St Kilda were similar (Fig. 1b).

Calanus copepods were also isotopically segregated among the geographic regions found for the breeder samples (MANOVA, Pillai, $F_{2,80} = 93.17$, P <0.0001, Fig. S2). Univariate analysis showed significant δ^{15} N differences between Iceland and the other regions (ANOVA, $F_{2,80} = 212.84$, P <0.0001; Scotland, t = 20.39, P <0.0001; Canada, t = 10.44, P

<0.0001), while Scotland and Canada were similar. $\delta^{13}C$ differences were found among all regions (ANOVA, $F_{2,80} = 208.55$, P <0.0001).

(B) Isotopic evidence for inter-colony movement using SIAR

Isotope values of pre-breeders were more variable than breeders, but (with the exception of a single outlier) pre-breeder values nested within breeder variation (Fig. 1b), suggesting good characterization of sources.

SIAR outputs revealed that of 134 pre-breeding Leach's storm-petrels caught at three colonies, many had isotopes dissimilar to their colony of capture (Fig. 2). For instance, 5 St Kildan pre-breeders had median probability estimates <25% for their colony of capture and were instead isotopically similar to Iceland (Fig. 2). Nineteen other pre-breeders (9 caught on St Kilda, 7 on Gull Island and 3 on Vestmannaeyjar) had mixed signatures (<50% for the colony of capture) indicating possible recent movement into their region of capture (Fig. 2). A comparison of model outputs (original vs "sensitivity") revealed changes to the input values for Gull Island had very little effect on the results (Table S2, Figs. S3 & S4).

(A) **DISCUSSION**

Using a novel application of stable isotope mixing models, we found evidence that prebreeding Leach's storm-petrels moved long-distances across the North Atlantic (spanning >3000km), among breeding colonies in Scotland, Iceland and Canada. These findings are significant for a number of reasons. First, they reveal the potential for long-distance dispersal at the ocean basin scale and lend credibility to the idea that Leach's storm-petrel colonies may act as a metapopulation. This provides important insight into population dynamics and also suggests this inter-colony exchange may have implications for buffering localised mortality (Bicknell *et al.*, 2012). Second, this work also provides evidence that the large pool of pre-breeders play a critical role in population and metapopulation dynamics of seabirds. We discuss the implications of these findings for population biology in general, and seabird conservation in particular, as well as the use of stable isotope mixing models to study movement of cryptic pre-breeders below.

(B) Stable isotopes and dispersal

The stable isotope mixing model used here obviates many of the constraints of studying intercolony connectivity of Leach's storm-petrels (e.g. small size, large population, exhaustive resighting effort). However the same technique could also be applied to many other taxa, as long as there is clear isotopic differentiation among subpopulations (regions). We found isotopic differences among breeders from the three colonies sampled across two years, and this difference was confirmed through analysis of zooplankton sampled in the same three regions in a single year (Fig. S1).

One assumption of mixing models is that consumer tissues are in equilibrium with their diet (Martínez del Rio & Wolf, 2005). Movements between isotopically distinct regions will not immediately be reflected in consumer tissues, for instance avian blood has a half-life of approximately 14 days (Bearhop *et al.*, 2002; Podlesak *et al.*, 2005). Therefore, the more time an individual spends in a new area the greater the isotopic similarity to that region as equilibrium approaches. Therefore the extent of movement among colonies assessed here is most likely to be an underestimate because only individuals moving relatively quickly between isotopically distinct regions will have values dissimilar to their colony of capture. Only 5 St Kilda pre-breeders can confidently be assessed as recent arrivals from the Icelandic region, but this still provides strong evidence for their movement between colonies and is

consistent with previous studies using other approaches (such as genetic markers) (Bicknell *et al.*, 2012).

Important to mixing model efficacy is good characterisation of potential sources - here the sources are the large foraging regions close to Leach's storm-petrel colonies, but may also include other at-sea areas such as over-winter locations and migratory pathways. Visual inspection of iso-space plots of sources and mixtures suggest we have achieved this (Fig. 1). However we did not sample all Leach's storm-petrel colonies in the North Atlantic but instead sampled the major oceanic water masses close to these colonies. For instance the vast majority of the North American breeding population are found in Canada (>95% (Huntingdon et al., 1996; Mitchell et al., 2004)), with the largest colonies on the north and south of Newfoundland. While we only sampled the north, given the large foraging range of this species (Hedd et al., 2009) they most likely integrate a signature derived from water masses around much of Newfoundland. Moreover, $\delta^{13}C$ and $\delta^{15}N$ values of copepods from a CPR route south of Newfoundland (Fig. S1), and data from the literature (Gulf of Maine (Estrada et al., 2005)), are similar to copepods from north Newfoundland suggesting our assumption is robust. A further possibility is that some Leach's storm-petrels sampled had recently returned from their wintering quarters or migratory staging grounds. While we cannot completely exclude this possibility, it seems unlikely given the close match of our source and mixture isotope values (Fig. 1). However, a single pre-breeder caught on St Kilda had an extremely high RBC δ^{13} C value (Fig. 1), which, because of large scale δ^{13} C gradients in the Atlantic (associated partly with temperature (Graham et al., 2010) and an inverse relationship with latitude (Paiva et al., 2010)) suggests this bird was probably a late arrival from lower latitude waters. We did not include this bird in further analyses, but it gives us some confidence that we can identify recently returning migrants in our analysis.

The CPR copepod samples confirmed regional isotopic differences but also revealed a different pattern of nitrogen and carbon isotope values at a lower trophic level (Fig. S2), compared to the breeding Leach's storm-petrel samples (Fig. 1b). The feeding ecology of Leach's storm-petrels is not known, or believed, to differ between regions (or between prebreeders and breeders), suggesting the change in isotopic pattern found at a higher trophic level may be attributed to food web structure/length in the regions. Further research and analyses (e.g. compound specific isotope analysis) is required to elucidate the potential cause of these differences, but highlights that variation among regions and across different trophic levels may be pronounced, which should be considered in future spatial or comparison studies.

Stable isotope mixing models provide an additional tool to infer movement of individuals between isotopically distinct regions or ecosystems. The use of additional elements in the analysis (e.g. oxygen or sulphur or highly-branched isoprenoids (Brown *et al.*, 2013)) may help provide more accuracy of the models through further differentiation of regions, but relies upon regionally distinct signatures. Combining stable isotope analyses with other indirect techniques, such as genetics, and/or direct methods, such as tracking, can build evidence for movement and dispersal of individuals. However for small and cryptic age-classes or species, such as Leach's storm-petrels, indirect methods and innovative analysis are currently the only realistic way to provide evidence for movement of individuals over large spatial scales.

(B) Predation on St Kilda

Our SIAR analysis suggests ~1 in 10 pre-breeders at St Kilda had recently arrived from another region, which could explain the continued persistence of this colony. The extremely

high, long-term predation by great skuas (~21,000 birds a year since 1996; Phillips *et al.*, 1999; Miles, 2010) would be unsustainable for this relatively small colony (~20,000 breeding pairs in 2006; Newson *et al.*, 2008). A rescue effect could manifest via compensatory recruitment (Votier *et al.*, 2008) or because skuas eat large numbers of prospecting prebreeders (Mougeot & Bretagnolle, 2000), or a combination of both. The lack of potential recent arrivals caught on Gull Island and Vestmannaeyjar compared with St Kilda, suggests pre-breeder movement among colonies may not be random, which would be indicative of a sink population sustained by larger source populations across the North Atlantic. Further work is required to determine whether this is the case or not.

Natal dispersal is thought to be important for inter-population exchange and metapopulation dynamics in a wide range of taxa (Paradis *et al.*, 1998; Clobert *et al.*, 2001; Inchausti & Weimerskirch, 2002; Telfer *et al.*, 2003; Berry *et al.*, 2004; Dittmann *et al.*, 2005; Caut *et al.*, 2008; Robinson & Beckerman, 2013), but can be very difficult to study. Here, we were able to provide indirect evidence of connectivity among populations through prospecting prebreeding individuals during the breeding period, and identify potential source/sink metapopulation dynamics over large spatial scales. Moreover this behaviour is likely helping the persistence of a local sub-population under extreme predation pressure. This further highlights the need to understand connectivity of disjunct populations of highly vagile species to be able to implement effective conservation measures at relevant spatial scales.

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(A) DATA ACCESSIBILITY

All the raw and corrected stable isotope data, model parameter files and model comparison results have been deposited in the Dryad repository; DOI: doi:10.5061/dryad.q2610

(A) **BIOSKETCH**

The authors have diverse research interests (e.g. dispersal, phylogeography, speciation, individual specialisation, foraging & spatial ecology, conservation genetics, conservation management and policy), which include the use of molecular techniques to address ecological

and evolutionary questions, and how the results may be applied to species and biodiversity conservation.

(A) AUTHOR CONTRIBUTIONS

AWJB, SCV, MEK, DTB and JBR conceived the ideas; AWJB, SCV, MC and JN collected and/or processed samples; AWJB and SCV analysed the data; AWJB and SCV led the writing.

Figure 1 a. Leach's storm-petrel colonies and CPR sampling locations in the North Atlantic. Black = Canadian region, Green = Icelandic region and Red = Scottish region. Stars = colony locations, Triangles = CPR sample locations. Arrows indicate direction and relative amount of pre-breeder movement inferred from SIAR results. b. δ^{13} C and δ^{15} N stable isotope values for breeders (bars; mean ± 2 sd) and individual pre-breeders (filled circles) by colony. Black = Gull Island, green = Vestmannaeyjar and red = St Kilda. (Photo taken by Richard Steel)



Figure 2 SIAR proportion estimates for individual pre-breeding Leach's storm-petrels caught at three colonies in the North Atlantic. **a**) Birds caught at Gull Island, Canada, **b**) Birds caught at St Kilda, Scotland, **c**) Birds caught at Vestmannaeyjar, Iceland. Black bar = Gull Island proportion estimate, red bar = St Kilda proportion estimate, green bar = Vestmannaeyjar proportion estimate.



SUPPORTING INFORMATION

Supporting Data file (

Table S1 {Leach's storm-petrel and *calanus* copepod sample sizes and tissue type. b = breeding adult Leach's storm-petrel, nb = non-breeding Leach's storm-petrel, * = each sample represents 10-20 individual copepods.}

Location	Species	Туре	Ν
<u>Colony</u> Gull Island, Canada Vestmannaeyjar, Iceland St Kilda, Scotland	Oceanodroma leucorhoa Oceanodroma leucorhoa Oceanodroma leucorhoa	Red Blood Cells Red Blood Cells Red Blood Cells	04 (b), 61 (nb) 12 (b), 22 (nb) 18 (b), 51 (nb)
<u>Atlantic Region</u> Newfoundland Shelf Reykjanes Ridge/Iceland Basin Shetland basin	Calanus finmarchicus Calanus finmarchicus Calanus helgolandicus	Whole organism Whole organism Whole organism	36* (2 CPR routes) 29* (2 CPR routes) 18* (1 CPR route)

Table S2 {Mean change in individual SIAR proportion estimates for sampled pre-breeding Leach's storm-petrels when comparing results between the original and sensitivity model (estimated Gull Island source δ^{13} C and δ^{15} N values and standard deviations).}

Pre-breeder colony		Change in individuals	Change in individuals
of capture	Model source	proportion estimates	proportion estimate
or capture		(range)	(mean ±1sd)
Gull Island, Canada	Gull Island St Kilda Vestmannaeyjar	-0.06 - 0.12 -0.04 - 0.05 -0.07 - 0.07	+0.01 ±0.06 -0.01 ±0.03 +0.01 ±0.04
St Kilda, Scotland	Gull Island St Kilda Vestmannaeyjar	-0.08 - 0.00 0.00 - 0.05 0.00 - 0.05	-0.05 ±0.03 +0.03 ±0.01 +0.03 ±0.02
Vestmannaeyjar , Iceland	Gull Island St Kilda Vestmannaeyjar	-0.080.01 0.00 - 0.04 0.01 - 0.07	-0.05 ±0.02 +0.02 ±0.01 +0.04 ±0.01



Figure S1 { δ^{13} C and δ^{15} N stable isotope values for breeders (bars; mean ± 2sd) and individual prebreeders (filled circles) used in the "sensitivity" model. Black = Gull Island (estimated values), green = Vestmannaeyjar and red = St Kilda.}



Figure S2 {*Calanus* copepod δ^{13} C and δ^{15} N stable isotope bivariate plot confirming the regional isotopic differences at this lower trophic level. Corrected means and 2 standard deviation bars for samples collected from the SAHFOS Continuous Plankton Recorder in oceanic regions off the coast of Newfoundland, Canada (both north and south), Iceland and Scotland (shown in Figure 1).}



Figure S3 {Sensitivity model results. SIAR median proportion estimates for individual pre-breeding Leach's storm-petrels caught at three colonies in the North Atlantic based on estimated Gull Island source δ^{13} C and δ^{15} N values and standard deviations. a) Birds caught at Gull Island, Canada, b) Birds caught at St Kilda, Scotland, c) Birds caught at Vestmannaeyjar, Iceland. Black bar = Gull Island proportion estimate, red bar = St Kilda proportion estimate, green bar = Vestmannaeyjar proportion estimate.}



Figure S4 {Mixing model proportion estimates and credibility intervals (25, 75 & 95%) for prebreeding Leach's storm-petrels caught at three colonies (Gull Island – Canada, St Kilda – Scotland and Vestmannaeyjar – Iceland). a. Model results by colony of capture (group) using original source δ^{13} C and δ^{15} N values and standard deviations, b. Sensitivity model results by colony of capture (group) using estimated Gull Island source δ^{13} C and δ^{15} N values and standard deviations.}