Natural geochemical markers reveal environmental history and population connectivity of common cuttlefish in the Atlantic Ocean and Mediterranean Sea

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Natural markers ($\delta^{13}C$ and $\delta^{18}O$ stable isotopes) in the cuttlebones of the European common cuttlefish (Sepia officinalis) were determined for individuals collected across a substantial portion of their range in the Northeast Atlantic Ocean (NEAO) and Mediterranean Sea. Cuttlebone $\delta^{13}C$ and $\delta^{18}O$ values were quantified for core and edge material to characterize geochemical signatures associated with early (juvenile) and recent (sub-adult/adult) life-history periods, respectively. Regional shifts in cuttlebone $\delta^{13}C$ and $\delta^{18}O$ values were detected across the 12 sites investigated. Individuals collected from sites in the NEAO displayed more enriched $\delta^{13}C$ and $\delta^{18}O$ values relative to sites in the Mediterranean Sea, with the latter also showing salient differences in both markers among western, central and eastern collection areas. Classification success based on cuttlebone $\delta^{13}C$ and $\delta^{18}O$ values to four geographical regions (NEAO, western, central and eastern Mediterranean Sea) was relatively high, suggesting that environmental conditions in each region were distinct and produced area-specific geochemical signatures on the cuttlebones of S. officinalis.

A modified $\delta^{13}C$ and $\delta^{18}O$ baseline was developed from sites proximal to the Strait of Gibraltar in both the NEAO and Mediterranean Sea to assess potential mixing through this corridor. Nearly, all (95%) of $\delta^{13}C$ and $\delta^{18}O$ signatures of S. officinalis collected in the area of the NEAO closest to the Strait of Gibraltar (Gulf of Cadiz) matched the signatures of specimens collected in the western Mediterranean, signifying potential movement and mixing of individuals through this passageway. This study extends the current application of these geochemical markers for assessing the natal origin and population connectivity of this species and potentially other taxa that inhabit this geographical area.
1. Background

The European common cuttlefish (*Sepia officinalis*) is an important fishery resource throughout the Northeast Atlantic Sea and adjacent waters of the Northeast Atlantic Ocean (NEAO) [1,2]. Due to large temporal shifts in the abundance and recruitment of *S. officinalis*, particularly for high-yield fisheries in certain areas (e.g. English Channel, Bay of Biscay [1]), there is renewed interest in their movement ecology and population connectivity [3]. This is primarily due to the fact that spatio-temporal shifts in distribution of marine organisms, particularly spawning stock biomass (SSB), can have profound implications for the management of harvested fisheries [4]. Current knowledge on the movement of *S. officinalis* is limited, preventing the implementation of conservation measures (e.g. time-area closures) for protecting highly suitable habitat or key areas used by spawning adults. Several different approaches have been used to clarify the movements and home range of *S. officinalis*, including conventional tagging [5], acoustic telemetry [6] and spatial shifts in catch from fishery-dependent data [7]. More recently, Dance et al. [8] used natural tracers in cuttlebones to investigate small-scale habitat shifts of this species in the Adriatic Sea. While earlier studies have shed important light on the habitat use and movement of *S. officinalis*, expanding the geographical scope of our knowledge on population connectivity is critical to their management.

Cuttlefish (class Cephalopoda) are characterized by the presence of an aragonitic internal shell (cuttlebone), and this gas-filled structure serves as a buoyancy control mechanism [9–11]. Similar to other biogenic carbonates found in cephalopods (statoliths) and fishes (otoliths), these aragonitic structures accrete material in a predictable manner and the microstructure retains information that records the timing of deposition [12]. In addition, the chemistry of material accreted in cuttlebones and other biogenic carbonates reflects ambient elemental or isotopic concentrations in seawater, and thus serve as natural markers for retrospective determination of past occurrences or environmental histories [8,13]. As a result, chemical markers in cuttlebones represent a powerful tool for predicting the natal origin, movement and population connectivity of cuttlefish [14].

The aim of the present study was to characterize regional variation in the geochemistry of cuttlebones of *S. officinalis* across a large section of its range in the NEAO and Mediterranean Sea. A common class of natural markers (stable C and O isotope ratios) was employed to determine whether distinct geochemical gradients exist within the Mediterranean Sea and areas outside this basin in the NEAO. We quantified δ¹³C and δ¹⁸O values for core and edge portions of the cuttlebone to contrast signatures and regional discrimination of habitats used during early (natal) and recent (time of collection) life-history periods, respectively. Multiple locations were sampled within each of the four major geographical regions investigated (NEAO and western, central, and eastern Mediterranean Sea), and regional discrimination (classification success) was determined for both cuttlebone core and edge signatures. We then used cuttlebone core (δ¹³C_CORE and δ¹⁸O_CORE) values to predict the natal origin of sub-adult/adult *S. officinalis* collected from a potential mixing area at the juncture between the NEAO and Mediterranean Sea. The spatial configuration of distinct geochemical provinces within areas investigated provides important insights into the suitability of these natural markers for addressing related ecological questions (ontogenetic habitat shifts) as well as population connectivity within the NEAO and Mediterranean Sea.

2. Methods

*Sepia officinalis* were collected between February 2013 and February 2014 across several locations in the NEAO and Mediterranean Sea, with samples falling into four major geographical regions: NEAO [Bay of Biscay to Gulf of Cadiz], western Mediterranean Sea, W-MED [Alboran Sea to Balearic Sea], central Mediterranean, C-MED [Ligurian Sea to Adriatic Sea], eastern Mediterranean, E-MED [Ionian Sea to Aegean Sea] (figure 1). Specimens were almost exclusively sub-adult/adult *S. officinalis* (table 1), and obtained from commercial trawlers, government run sampling.
Table 1. Summary of *Sepia officinalis* collected in the Northeast Atlantic Ocean and Mediterranean Sea with information on site code (A-L), location, collection date, sample number (N) and mean mantle length (ML) of specimens from each site. ◊ Collection area in Bay of Cadiz outside Strait of Gibraltar used to assess exchange between the Northeast Atlantic Ocean and Mediterranean Sea.

| site  | location or sampling port | collection date (MO/HR) | N       | mean ML ±1 s.d. (mm) |
|-------|---------------------------|-------------------------|---------|---------------------|
| A     | La Rochelle, France       | June 2013               | 20      | 143.7 ±9.4          |
| B     | San Sebastian, Spain      | February–June 2013      | 44      | 121.0 ±11.3         |
| C     | Cascais, Portugal         | June 2013               | 24      | 79.2 ±36.9          |
| D     | Olhão, Portugal           | June 2013               | 19      | 79.8 ±16.3          |
| ◊     | Cadiz, Spain              | June 2013               | 20      | 110.0 ±18.9         |
| E     | Marbella, Spain           | June 2013               | 10      | 111.4 ±20.3         |
| F     | Cubellas, Spain           | June 2013               | 19      | 91.8 ±7.8           |
| G     | Sardinia, Italy           | September–October 2013  | 20      | 69.0 ±21.7          |
| H     | La Spezia, Italy          | October 2013            | 22      | 60.5 ±9.0           |
| I     | Venice, Italy             | October 2013            | 17      | 52.1 ±4.9           |
| J     | Galaxidi, Greece          | July–September 2013     | 20      | 103.0 ±16.9         |
| K     | Athens, Greece            | February 2014           | 20      | 88.3 ±16.5          |
| L     | Santorini, Greece         | February 2014           | 6       | 98.3 ±12.4          |

*DDenoted on figure 1.*

programmes, artisanal fishers, and local fish markets. Cuttlebones were extracted, cleaned and dried in the laboratory. Core and edge material for δ¹³C and δ¹⁸O analysis was collected using the protocol described previously [8]. Briefly, approximately 1-cm sections (approx. 1–3+ month interval [15]) of material were isolated from the posterior origin (core) and anterior margin (edge) of each cuttlebone (figure 2). For each section, material was carefully removed from the phragmocone portion of the cuttlebone by sampling from the ventral surface upward to the hypostracum (ventral layer of the dorsal shield). Material was isolated from each cuttlebone with a scalpel and powdered.

Cuttlebone δ¹³C and δ¹⁸O were determined using an automated carbonate preparation device (KIEL-III Thermo Fisher Scientific, Inc.) coupled to an isotope ratio monitoring mass spectrometer (Finnigan MAT 252 Thermo Fisher Scientific, Inc.) at the University of Arizona Environmental Isotope Laboratory. Powdered cuttlebone samples were reacted with dehydrated phosphoric acid under vacuum at 70°C. The isotope ratio measurement was calibrated based on repeated measurements of the National Bureau of Standards (NBS), NBS-19 and NBS-18, with six standards run for every 40 samples; precision was approximately 0.08‰ and 0.11‰ for δ¹³C and δ¹⁸O, respectively. Cuttlebone δ¹³C and δ¹⁸O are reported relative to the Vienna Pee Dee Belemnite (VPDB) scale after comparison to an in-house laboratory standard calibrated to VPDB.

Multivariate analysis of variance (MANOVA) was used to test for regional difference in cuttlebone core and edge δ¹³C and δ¹⁸O. In addition, univariate contrasts (ANOVA) were performed on individual markers to assist in the identification of the most influential markers for regional discriminations. MANOVA significance used Pillai’s trace statistic, and significance for all ANOVA and MANOVA tests was based on an alpha level of 0.05. Results from ANOVA tests were examined further with Tukey’s honestly significant difference (HSD) test to determine whether group means between specific regions were different. Paired *t*-tests were also performed between core and edge values for both δ¹³C and δ¹⁸O to determine whether early (core) and recent (edge) life-history signatures from the same individual differed. Canonical discriminant analysis (CDA) was used to display multivariate means of cuttlebone δ¹³C and δ¹⁸O, and discriminant function coefficients were incorporated into CDA plots as vectors from a grand mean to show the discriminatory influence of δ¹³C and δ¹⁸O by region. Quadratic discriminant function analysis (QDFA) was then used to determine the classification accuracy of cuttlebone δ¹³C and δ¹⁸O signatures for assigning individuals to their respective region (i.e. collection area). We selected QDFA over other multivariate approaches to assess classification success based on natural makers because it is preferred when the variance–covariance matrix of our predictor variables differs. Also, QDFA does not have the assumption of homogeneity of covariance matrices and is robust to moderate deviations from normality [16].

Natal origin of *S. officinalis* collected near the Strait of Gibraltar in the NEAO was predicted using direct maximum-likelihood estimates (MLE) of stock composition [17,18]. Mixed-stock analysis was run under bootstrap mode to obtain standard deviations around estimated proportions (error terms) with 500 simulations [19]. For our assessment of NEAO and W-MED mixing, we limited our source populations to the two nearest collection sites on each side of the Strait of Gibraltar (NEAO = sites C, D; W-MED = sites E, F; figure 1), and our unknown population was based on larger (greater than 100 mm mantle length) *S. officinalis* collected proximal to the Strait of Gibraltar in the Bay of Cadiz (distance approx. 80 km). Cuttlebone δ¹³C and δ¹⁸O values of *S. officinalis* collected in the Bay of Cadiz were all within 95% confidence ellipses of individuals from the two general regions used to create the baseline and thus we assumed that bias due to the presence of individuals from nurseries not included or not sampled was minimal.
3. Results

3.1. Regional variation in cuttlebone $\delta^{13}C$ and $\delta^{18}O$

Significant spatial differences in cuttlebone $\delta^{13}C_{CORE}$ and $\delta^{18}O_{CORE}$ signatures (early life history) were detected for $S.\ officinalis$ among the collection sites (MANOVA, $p < 0.01$; figure 3). Mean cuttlebone $\delta^{13}C_{CORE}$ values ranged from $-2.69\%$ to $-5.10\%$ across the 12 sites, and a distinct trend was observed for $S.\ officinalis$ from sampling sites in the NEAO versus sites within the Mediterranean Sea. Individuals collected in the NEAO generally displayed more enriched cuttlebone $\delta^{13}C_{CORE}$ values (mean ± 1 SD: $-3.16\% \pm 0.67\%$) relative to all three regions in the Mediterranean Sea: W-MED ($-3.94\% \pm 0.78\%$), C-MED ($-4.65\% \pm 0.95\%$) and E-MED ($-3.97\% \pm 0.78\%$). Univariate contrasts indicated that cuttlebone $\delta^{13}C_{CORE}$ values were significantly different among the four regions (ANOVA, $p < 0.01$), and Tukey HSD test indicated that $\delta^{13}C_{CORE}$ values for $S.\ officinalis$ from the NEAO region were significantly different from all three regions within the Mediterranean Sea. Site-specific variation in cuttlebone $\delta^{18}O_{CORE}$ was also present and mean values ranged from $-0.2\%$ to $1.3\%$ across the 12 sites sampled. Similar to $\delta^{13}C_{CORE}$, cuttlebone $\delta^{18}O_{CORE}$ values were significantly different among the four geographical regions (ANOVA, $p < 0.01$), and mean values were higher in the NEAO ($0.92\% \pm 0.44\%$) compared to sites within the Mediterranean Sea: W-MED ($0.53\% \pm 0.37\%$), C-MED ($0.04\% \pm 0.95\%$) and E-MED ($0.62\% \pm 0.67\%$). Again, Tukey HSD tests indicated that $\delta^{18}O_{CORE}$ values for individuals collected in the NEAO region were significantly different from all three regions of the Mediterranean Sea.

Similarly, spatial variation in cuttlebone $\delta^{13}C_{EDGE}$ and $\delta^{18}O_{EDGE}$ values (recent life history) were detected for $S.\ officinalis$ (MANOVA, $p < 0.01$; figure 4). Mean cuttlebone $\delta^{13}C_{EDGE}$ values ranged from $-3.49\%$ to $-0.02\%$ across the 12 sites sampled. Again, a discernible NEAO versus Mediterranean Sea pattern was observed for cuttlebone $\delta^{13}C_{EDGE}$ with mean values significantly higher in the NEAO ($-0.73\% \pm 1.10\%$) relative to the three regions within the Mediterranean Sea: W-MED ($-2.70\% \pm 0.87\%$), C-MED ($-2.59\% \pm 0.80\%$), E-MED ($-2.21\% \pm 1.60\%$) (ANOVA, $p < 0.01$, Tukey HSD $p < 0.05$). Spatial variation in cuttlebone $\delta^{18}O_{EDGE}$ was also pronounced across the 12 sites (range: $0.50\%$ to $2.42\%$). Although a significant region effect was observed (ANOVA, $p < 0.01$), $S.\ officinalis$ sampled from the NEAO and two regions within the Mediterranean Sea displayed similar cuttlebone $\delta^{18}O_{EDGE}$ values: NEAO ($1.88\% \pm 0.54\%$), W-MED ($1.64\% \pm 0.37\%$) and E-MED ($1.91\% \pm 0.63\%$) (Tukey HSD $p > 0.05$). Region-specific variation in cuttlebone $\delta^{18}O_{EDGE}$ was due to $S.\ officinalis$ sampled from the C-MED having significantly lower $\delta^{18}O_{EDGE}$ values ($0.93\% \pm 0.55\%$) relative to the other three regions (Tukey HSD $p < 0.05$).

Pair comparisons of cuttlebone core and edge material were also assessed to determine whether ontogenetic changes in habitat or environmental conditions may have occurred between early and recent life-history periods. Although general regional trends persisted for both cuttlebone core and edge samples, differences were detected between the two sections of the cuttlebone assayed for both markers. Cuttlebone $\delta^{13}C_{CORE}$ and $\delta^{13}C_{EDGE}$ values were significantly different (paired $t$-test, $p < 0.01$), and mean cuttlebone $\delta^{13}C_{EDGE}$ values were approximately $2\%$ higher than $\delta^{13}C_{CORE}$ values (mean difference ± 1 s.d.: $2.03\% \pm 1.48\%$). Observed differences were highly variable among $S.\ officinalis$ from the 12 sampling sites (figures 3 and 4). Similarly, differences between the early and recent life-history periods were observed for $\delta^{18}O$ and cuttlebone $\delta^{18}O_{EDGE}$ values were significantly higher than $\delta^{18}O_{CORE}$ values (mean difference $1.00\% \pm 0.93\%$).

3.2. Classification to natal sites and stock discrimination

Canonical discriminant analysis based on cuttlebone $\delta^{13}C$ and $\delta^{18}O$ of $S.\ officinalis$ showed clear separation of 95% confidence ellipses for each centroid based on cuttlebone core (figure 5a) and edge (figure 5b) values. Biplot vectors from a grand mean indicated that discrimination to the four regions was influenced.
by both $\delta^{13}$C and $\delta^{18}$O. QDFA based on cuttlebone $\delta^{13}$C$_{\text{CORE}}$ and $\delta^{18}$O$_{\text{CORE}}$ (early life history) of $S. officinalis$ resulted in cross-validated classification success of 62.1% to the four geographical regions. QDFA parameterized with $\delta^{13}$C$_{\text{EDGE}}$ and $\delta^{18}$O$_{\text{EDGE}}$ (recent life history) showed a slight reduction in classification success to 60.2%. Combining cuttlebone material from the two life-history periods (core and edge) resulted in the highest cross-validated classification success (75.4%) to the four regions, indicating that regional-specific chemical differences in cuttlebone $\delta^{13}$C and $\delta^{18}$O at both life-history periods were useful for discriminating $S. officinalis$ from the geographical areas sampled. We also ran QDFA on a subset of sites in both the NEAO (site C, D) and W-MED (sites E, F) to evaluate the suitability of using $\delta^{13}$C$_{\text{CORE}}$ and $\delta^{18}$O$_{\text{CORE}}$ signatures for assessing population exchange, and classification success to these two regions was 84.5%.

### 3.3. Population connectivity between NEAO and Mediterranean Sea

Predictions of natal origin for $S. officinalis$ collected in close proximity to the Strait of Gibraltar (Gulf of Cadiz) were used to assess potential exchange (i.e. movement) of individuals between the NEAO and Mediterranean Sea. A modified baseline comprised only of cuttlebone $\delta^{13}$C$_{\text{CORE}}$ and $\delta^{18}$O$_{\text{CORE}}$ values for individuals collected from two sites closest to the
Strait of Gibraltar in both the NEAO and Mediterranean Sea (NEAO = C, D; W-MED = E, F) was used to predict the natal origin of *S. officinalis* from the Gulf of Cadiz. The majority of δ¹³C_CORE and δ¹⁸O_CORE values of *S. officinalis* collected in the Gulf of Cadiz matched bivariate kernel density estimates of cuttlebone δ¹³C and δ¹⁸O for individuals from the W-MED (E, F) rather than NEAO (C, D) (figure 6). MLE-based predictions of natal origin using the modified baseline indicated that *S. officinalis* in our Gulf of Cadiz sample were almost exclusively of western Mediterranean origin (95.4% ± 8.2%).

### 4. Discussion

Spatial variation in cuttlebone δ¹³C and δ¹⁸O of *S. officinalis* was observed across the large geographical scale investigated, with salient regional-specific differences in geochemical signatures present for both early (core) and recent (edge) life-history periods detected. Classification to the four regions (NEAO, W-MED, C-MED, E-MED) based on mixed-stock models parameterized with cuttlebone δ¹³C and δ¹⁸O was also similar using core or edge signatures. Regional discrimination based on these natural markers indicated that environmental conditions and drivers of seawater and/or cuttlebone geochemistry likely function at broad spatial scales in the NEAO and Mediterranean Sea. The four-region framework adopted for our study appears to adequately capture major geographical trends in cuttlebone geochemistry at spatial scales that are often isotopically distinct within marginal seas [20, 21]. Noticeable trends in cuttlebone δ¹³C of *S. officinalis* were present among sites and regions, although site to site variability within most regions was relatively modest, particularly for δ¹³C_CORE. A notable trend included cuttlebone δ¹⁰C_CORE values being more positive for individuals collected from nearly all sites in the NEAO (A-D) relative to sites within the Mediterranean Sea (E-L). We also observed a gradient of decreasing δ¹⁰C_CORE values from the NEAO to the C-MED, followed by a slight increase at sites in the E-MED. Similar to our findings, conspicuous differences in phytoplankton δ¹³C (δ¹³C PHY) and zooplankton δ¹³C (δ¹³C_ZOO) have been reported between areas of the NEAO and Mediterranean Sea where *S. officinalis* were collected, with δ¹³C values more positive in the NEAO relative to the Mediterranean Sea [21]. This may be attributed to differences in primary productivity and phytoplankton growth rates that are reported to be higher in areas sampled in the NEAO, possibly leading to higher δ¹³C PHY in this region [22]. In addition, it is possible that cuttlebone δ¹³C_CORE values may be influenced to some degree by maternally derived geochemical signatures, which can be conserved in the cores of biogenic hard parts [23]. We also observed generally higher δ¹³C_CORE and δ¹⁰C_EDGE values for *S. officinalis* collected in the E-MED relative to the C-MED and W-MED, and this appears due to higher water temperatures (increases phytoplankton growth rate) and elevated salinity in the E-MED; both parameters are positively related to δ¹³C PHY [24]. Finally, it is important to note that sites within each region, particularly for δ¹³CEDGE, generally maintained similar mean values, particularly in the NEAO, C-MED and E-MED. This is noteworthy because sites within each respective region were often separated by considerable distances (greater than 500 km) and in different marginal seas (e.g. Tyrrhenian and Adriatic in C-MED), suggesting that the resolution of this marker appears to be functional at the geographical scale investigated (i.e. four regions).

Regional distinctions in cuttlebone δ¹⁸O_CORE and δ¹⁸O_EDGE values between sites in the NEAO and Mediterranean Sea were also detected, and it is assumed that δ¹⁸O in biogenic carbonates reflects seawater δ¹⁸O(SEAWATER) values at the ocean-basin scale [25]. Patterns in cuttlebone δ¹⁸O for *S. officinalis* did not entirely match previously reported gradients in δ¹⁸O(SEAWATER) for offshore waters in the NEAO and Mediterranean Sea [20, 26], suggesting that other physico-chemical or biological factors likely play a role in determining cuttlebone δ¹⁸O. The mismatch between δ¹⁸O(SEAWATER) and cuttlebone δ¹⁸O is possibly due in part to *S. officinalis* occupying inshore/near shore waters during early life [3], and such areas are often heavily influenced by freshwater inflow which alter δ¹⁸O values. Therefore, δ¹⁸O(SEAWATER) values in near shore nursery habitats of *S. officinalis* may not align well with predictions from ocean models used to predict geographical variation in δ¹⁸O(SEAWATER). The most noticeable discontinuity in cuttlebone δ¹⁸O_Core occurred between sites in the NEAO and Mediterranean Sea. Also, both cuttlebone δ¹⁸O_CORE and δ¹⁸O_EDGE values for sites in the E-MED off Greece were generally higher compared to all other sites in the Mediterranean Sea, which is not unexpected because δ¹⁸O(SEAWATER) is predicted to be highest in the eastern part of this marginal sea [20]. Elevated δ¹⁸O(SEAWATER) and cuttlebone δ¹⁸O_CORE and δ¹⁸O_EDGE in the E-MED is assumed to be due to increased salinity (evaporation), which causes a disproportionate removal of the lighter isotope (¹⁸O) in water vapour, leaving behind more of the heavier isotope and resulting in higher overall δ¹⁸O(SEAWATER) [27].

Classification success to the four regions sampled using cuttlebone δ¹³C and δ¹⁸O for early (core) and recent (edge) life-history periods indicated that *S. officinalis* collected from the NEAO, W-MED, C-MED and E-MED experienced different environmental conditions. We also observed that cuttlebone δ¹³C and δ¹⁸O values at sites within a region were often statistically similar, supporting the delineations used in our regional framework. This also indicates that environmental conditions experienced by *S. officinalis* were distinctive at the regional scale, and possibly associated with population structure. A recent study using genetic markers reported that populations

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**Figure 6.** Contour plots of cuttlebone δ¹³C and δ¹⁸O values for Sepia officinalis for the two locations nearest to the Strait of Gibraltar in both the NEAO (sites C and D) and W-MED (sites E and F) was used for prediction of natal origin. Bivariate kernel density estimated at four levels for samples from both NEAO and W-MED (25%, 50%, 75% and 100%). Cuttlebone δ¹³C and δ¹⁸O values for *S. officinalis* of unknown origin collected near the Strait of Gibraltar are shown with X symbol.
of *S. officinalis* in the Mediterranean Sea are fragmented into several subpopulations [28], and the geographical areas of their genetic clusters were remarkably similar to our regional resolution using geochemical markers. Interestingly, all of our sampling sites in both the C-MED (G, H, I) and E-MED (J, K, L) were each associated with unique genetic clusters (i.e. subpopulations). Similarly, moving into the western part of the basin and into the NEAO, the population structure again from genetics was similar to results from cuttlebone isotope geochemistry. Genetic data indicated the presence of two genetic clusters in the NEAO with a break between our collection sites in the Bay of Biscay (A, B) and sites near the Strait of Gibraltar (C, D). Similarly, we detected a shift in both cuttlebone δ18O_CORE and δ18O_EDGE between these two geographical areas (A, B versus C, D) (figures 4b and 5b). Genetic data also showed that sites on both sides of the Strait of Gibraltar (NEAO = C, D; W-MED = E) were part of the same subpopulation cluster, indicating that gene flow occurs through this corridor [28].

Application of these geochemical markers was also used to investigate exchange of *S. officinalis* between the NEAO and Mediterranean Sea over a restricted geographical scope. Previous mark–recapture and electronic tagging studies suggest that based on observed rates of movement (km day−1) *S. officinalis* are easily capable of travelling several hundred kilometres [3,5]. Therefore, a modified baseline was developed using a subset of sites proximal to the Strait of Gibraltar in both the NEAO and W-MED to assess potential population connectivity. Mixed-stock predictions of natal origin showed substantial exchange with individuals of W-MED origin presumably moving through the Strait of Gibraltar and into the NEAO. Migrations from natal sites in the Mediterranean Sea to areas in the NEAO occur for highly migratory species such as European eel Anguilla anguilla [29] and Atlantic bluefin tuna Thunnus thynnus [30]; however, connectivity between the NEAO and Mediterranean Sea for species assumed to display limited movements is ostensibly uncommon, and this type of egress activity has not been previously reported for *S. officinalis*. Nevertheless, our observation of *S. officinalis* of W-MED origin being present in the NEAO proximal to the Strait of Gibraltar is in accord with findings on stock structure using genetic markers, with Drábková et al. [28] reporting that *S. officinalis* from the NEAO and W-MED on each side of this passageway were grouped into the same subpopulation (genetic cluster). Because *S. officinalis* lays egg clusters that are typically attached to benthic plants or substrate and hatch fully developed [31], their dispersal potential relative to species with pelagic larvae is likely very limited. Consequently, gene flow through the Strait of Gibraltar requires active movement of individuals that is likely facilitated by strong surface currents flowing through this corridor [32] and offshore overwintering movements [3], which supports our finding of W-MED origin recruits west of this corridor in the NEAO. Still, it is important to note that our sample was limited in size and therefore may not accurately represent the stock composition of W-MED and NEAO recruits in the Bay of Cadiz.

Finally, higher cuttlebone δ13C and δ18O in edge relative to core material is indicative of the onontogenic shifts that typically occurs for *S. officinalis*, with juveniles often inhabiting bays, estuaries, or near shore environments [31]. Physico-chemical conditions in these nurseries (e.g. lower salinity and higher temperature) are influenced by near shore processes (e.g. freshwater inflow), which appear responsible for individuals displaying distinct δ13C and δ18O signatures. Increases in cuttlebone δ13C_EDGE and δ18O_EDGE at most of the sites surveyed is in accord with the shift of larger/older individuals to deeper, cooler and more saline offshore waters [8]. Increases in δ18O in the biogenic structures of bony fishes [33,34] and sharks [35] have also been attributed to inshore–offshore transitions or shifts in salinity exposure due to the positive relationship between salinity and δ18O_SEAWATER [36] and the inverse relationship of δ18O with temperature [8], which supports the cuttlebone core to edge change observed here for *S. officinalis*. Although ontogenetic shifts in cuttlebone δ13C and δ18O appear to correspond to inshore–offshore migrations, a fair degree of natural variability was present between core and edge signatures both within and across sites. Our finding is not surprising and due in part to the fact that physico-chemical differences between inshore–offshore waters can vary widely across broad geographical ranges [37]. It is also important to note that variability in both markers was more pronounced for cuttlebone edge material. This was likely due in part to the variation in the collection month and size/age composition while core material reliably characterized the early life period (nursery habitat) of all *S. officinalis* in our sample.

5. Conclusion

Our findings support the use of cuttlebone geochemistry to gain further insight into past environmental conditions and population connectivity of *S. officinalis*. Geographical variation in cuttlebone δ13C and δ18O implies that physico-chemical conditions experienced by *S. officinalis* during both early life and later periods in life are different among the four regions investigated. Given that environmental conditions are known to influence local adaptations (i.e. adaptive response) and population genetics [38], region-specific differences in environmental conditions experienced by *S. officinalis* may serve as a selective force leading to population structure (subpopulations) in the NEAO and Mediterranean Sea. Region-specific variation in cuttlebone δ13C and δ18O also provides additional insight regarding the spatial resolution and suitability of these geochemical markers for future investigations of the population connectivity of *S. officinalis* as well as highly migratory taxa with home ranges that include areas within and outside the Mediterranean Sea (e.g. Atlantic bluefin tuna; Rooker & Secor [39]). Although regional-scale variation in cuttlebone δ13C and δ18O was evident, we also demonstrate that the approach shows promise for applications at smaller spatial scales (site-specific) and reveal for the first time that *S. officinalis* with putative Mediterranean Sea (W-MED) signatures were observed in the NEAO, suggesting that individuals of Mediterranean origin may actively move through the Strait of Gibraltar and enter the NEAO.

Data accessibility. Data are available as part of the electronic supplementary material.

Authors’ contributions. J.R.R. conducted data analysis and wrote the main manuscript text; J.R.R. and R.J.D.W. designed the study; J.R.R., R.J.D.W., M.A.D., and J.L. processed samples; J.R.R., R.J.D.W., P.A., H.A., M.B., G.B., I.F., T.L., P.M., R.R., I.S., A.V.S. and R.V.; all authors provided comments on the manuscript.

Competing interests. We declare we have no competing interests.

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