Deep Sea Research Part I: Oceanographic Research Papers
April 2022, Volume 182, Pages 103708 (11p.)

https://archimer.ifremer.fr

https://doi.org/10.1016/j.dsr.2022.103708 https://archimer.ifremer.fr/doc/00750/86170/

Diet consistency but large-scale isotopic variations in a deep-sea shark: The case of the velvet belly lantern shark, *Etmopterus spinax*, in the northeastern Atlantic region and Mediterranean Sea

Besnard Lucien ^{1,*}, Duchatelet Laurent ², Bird Christopher S. ³, Le Croizier Gaël ⁴, Michel Loic ⁵, Pinte Nicolas ², Lepoint Gilles ⁶, Schaal Gauthier ¹, Vieira Rui P. ³, Gonçalves Jorge M.S. ⁷, Martin Ulrich ², Mallefet Jérôme ²

- ¹ Univ Brest, CNRS, IRD, Ifremer, LEMAR, F-29280, Plouzane, France
- ² University of Louvain UCLouvain, Earth and Life Institute, Marine Biology Laboratory, Croix du Sud 3, 1348, Louvain-la-Neuve, Belgium
- ³ Centre for Environment, Fisheries, & Aquaculture Sciences (CEFAS), Pakefield Road, Lowestoft, UK
- ⁴ UMR Géosciences Environnement Toulouse (GET), Observatoire Midi Pyrénées (OMP), 14 Avenue Edouard Belin, 31400, Toulouse, France
- ⁵ Univ Brest, Ifremer, CNRS, Unité Biologie et Ecologie des Ecosystèmes marins Profonds (BEEP), F-29280, Plouzane, France
- ⁶ University of Liège (ULg), Laboratoire d'Ecologie trophique et isotopique (LETIS), B6 Sart Tilman, 4000, Liège, Belgium
- ⁷ CCMAR Centro de Ciências do Mar, Universidade do Algarve, 8005-139, Faro, Portugal
- * Corresponding author: Lucien Besnard, email address: lbesnard.research@gmail.com

Abstract:

Deep-sea elasmobranchs are commonly reported as bycatch of deep-sea fisheries and their subsequent loss has been highlighted as a long-running concern to the ecosystem ecological functioning. To understand the possible consequences of their removal, information on basic ecological traits, such as diet and foraging strategies, is needed. Such aspects have been widely studied through stomach content analysis but the lack of long-term dietary information requires other tools to be used such as stable isotopes. This study examines nitrogen and carbon isotope compositions of the velvet belly lantern shark, Etmopterus spinax, one of the most impacted shark species in Northeastern Atlantic fisheries as a result of accidental catches. E. spinax was sampled at four different locations, characterized by contrasting oceanographic and ecological conditions: the western Mediterranean Sea (near the Balearic Islands), the southern Iberian upwelling system, Rockall Trough and southwestern Norwegian fjords. Stomach content analysis revealed similar prey species among sites, with a diet dominated by Euphausiacea (mostly Meganyctiphanes norvegica) and an ontogenetic shift towards small teleost fishes, cephalopods or other crustaceans. Despite these similarities, muscle stable isotope compositions differed across sampled locations. Rather than clear dietary differences, the contrasted isotopic values are likely to reflect differences in environmental settings and biogeochemical processes affecting nutrient dynamics at the base of the food webs.

Highlights

► *E. spinax* were sampled across the northeastern Atlantic and Mediterranean Sea. ► Stomach contents suggested a similar diet. ► *E. spinax* fed mainly on Euphausiacea with an ontogenetic diet shift. ► Muscle carbon and nitrogen isotope compositions differed between locations. ► Isotopic differences probably resulted from mechanisms affecting the baselines.

Keywords: Trophic ecology, Stable Isotopes, Stomach content, Mesopredator, Food webs, Benthopelagic predator.

1. Introduction

Under the lootprint of anthropogenic activities and climate change, many coastal and
epipelagic fisheries have significantly declined (Chavez et al., 2003; Pinsky et al., 2011; Tu et
al., 2018), diverting fishing efforts toward deep-sea stocks (Bailey et al., 2009; Devine et al.,
2006; Priede et al., 2011; Vieira et al., 2019). As a consequence, significant population
decreases have been observed in these deep water ecosystems with unknown and potentially
disruptive impacts on ecological processes and functions (Benn et al., 2010; Vieira et al., 2020).
However, despite their ecological importance, fragility and current state of exploitation, deep-
sea ecosystems are still relatively understudied compared to their shallower counterparts
(Benn et al., 2010; Thurber et al., 2014). In order to predict how deep-sea ecosystems will
respond to natural or human-induced changes, information on the functioning of deep-sea
communities is urgently needed (Howell et al., 2021).
Previous descriptions of deep-sea habitats have established that food webs are complexly
structured and include a range of trophic levels fuelled by a mixture of primary production,
secondary production and benthic recycling (Newman et al., 2011; Shipley et al., 2017b;
Trueman et al., 2014). Due to the absence of light for autochthonous primary production,
deep-sea fauna relies on the downward vertical transport of nutrients (Polunin et al., 2001;
Preciado et al., 2017), either actively by species diel vertical migrations or passively due to
particulate organic matter sinking and re-suspension (Trueman et al., 2014). Within these food
webs, deep-sea elasmobranch species are meso- to top predators (Churchill et al., 2015;
Simpfendorfer and Kyne, 2009) which are of critical importance through their potential top-
down or more complex regulation of communities (Heithaus et al., 2008; Shipley et al., 2017a).
By connecting different depth layers, they also influence energy flux and carbon cycling,
further enhancing their key ecological role (Trueman et al., 2014). Unfortunately, information

46 on deep-sea sharks is still lacking due to the logistical challenges of accessing and studying 47 them (Hussey et al., 2018; Moura et al., 2018; Simpfendorfer and Kyne, 2009). 48 In deep-sea sharks, most trophic studies have relied upon stomach content analysis (Barría et al., 2018; Cortés, 1999). Even if this approach allows a complete qualitative description of the 49 diet, it requires a large number of samples, represents only a snapshot of the last meal(s) and 50 51 differences in digestion rates may also bias the importance of prey items (Albo-Puigserver et al., 2015). In contrast, biochemical tracers such as stable isotope ratios of carbon (δ^{13} C) and 52 nitrogen (δ^{15} N) can help to describe the trophic structure, niche width and energy fluxes on a 53 54 long-term basis with potential quantitative approaches (Layman et al., 2012; Shipley et al., 2017a). Carbon isotope signatures are considered as a good proxy to characterize the primary 55 producers at the base of food webs (Fry and Sherr, 1984; Layman et al., 2012). Nitrogen 56 isotope composition is mainly used as a proxy of trophic position due to a global increase in 57 58 δ¹⁵N signal from prey to predator (Cabana and Rasmussen, 1994; Post, 2002) and has been linked to foraging depth in benthopelagic communities (Trueman et al., 2014). By extension, 59 60 the combination of carbon and nitrogen isotopes constitutes a proxy of the trophic niche crucial for assessing the ecological role of a given species or population (Newsome et al., 61 2007). 62 Deep-sea shark isotopic values depend on their diet but also on the isotopic compositions at 63 64 the base of food webs, which exhibit spatial and temporal variations (Magozzi et al., 2017; Somes et al., 2010). Environmental conditions and local biogeochemical processes are known 65 to affect baseline isotopic profiles due to changes in nutrient dynamics. For example, 66 latitudinal differences in baseline isotopic values can be influenced by temperature (Magozzi 67 et al., 2017; Rau et al., 1997), river discharges (Chouvelon et al., 2012) and upwelling events 68

69 (Lopez-Lopez et al., 2017; Puccinelli et al., 2019). The extent to which large-scale spatial variations in isotopic baselines are reflected in deep-sea ichtyofauna remains unclear. 70 Moreover, deep-sea sharks are mobile species that can shift their feeding ground while 71 72 migrating. Isotopically, this shift will follow baseline changes and these variations need to be considered when analysing the trophic niche of a species over time or space (Bird et al., 2018; 73 74 Lorrain et al., 2015). 75 Along the northeastern Atlantic and Mediterranean continental shelfs and slopes, the trophic 76 ecology of a small deep-sea shark, the velvet belly lantern shark, Etmopterus spinax (Linnaeus, 77 1758), has been extensively studied using stomach content analysis (Klimpel et al., 2003; Neiva et al., 2006; Valls et al., 2017). Beyond those habitats, luminous velvet belly lantern sharks 78 79 inhabit deep layers of fjords in Norway (Claes et al., 2010; Duchatelet et al., 2021). The first objective of this study is to present a description of the species diet inside these semi-enclosed 80 81 habitats. More generally, throughout its distribution area, only a few studies took into account stable isotopes to specifically investigate E. spinax trophic habitat (Albo-Puigserver et al., 82 2015; Valls et al., 2017). Consequently, the second objective of this study is to investigate 83 muscle carbon and nitrogen isotopic compositions of E. spinax at four different locations in 84 85 the Mediterranean Sea and in the northeastern Atlantic to gain long-term dietary information 86 on this species. Information on the trophic ecology of E. spinax gathered by stomach content 87 analysis were ultimately compared to stable isotope values and discussed in terms of local habitat ecological characteristics and physical processes affecting nutrient dynamics. 88

2. MATERIALS AND METHODS

89

90

2.1. Stomach content

In Norway, stomach content analysis of *Etmopterus spinax* came from eleven scientific expeditions from 2008 to 2012 inside the Raunefjord (Figure 1). Based on morphological analysis of less-digested or undigested component, preys were identified at the lowest taxonomic level possible. For Euphausiacea, eyes were digested at a lower rate than the rest of the body parts. Therefore, each pair of eyes was identified as one Euphausiacea individual unless morphological characteristics allowed identification to the species level. Cephalopods were identified through their remaining beaks. For large fishes, species identification was possible when the whole body was present or based on dental bone dimension. To assess and compare prey composition, four indices were calculated: the numeric percentage %N (a prey item abundance as a percent of the total prey abundance), the gravimetric percentage %W (a prey item remaining mass as a percent of the total prey mass), the occurrence percentage %O (number of stomachs containing a prey item in percent compare to all stomachs) and finally the index of relative importance IRI: IRI = (%N + %W) × %O (Cortés, 1997). Empty stomachs were not considered in indices calculation.

2.2. Stable isotope analysis

From 2014 to 2017, Etmopterus spinax specimens were sampled in four different areas of the northeast Atlantic and the Mediterranean Sea (Figure 1). Individuals from Rockall Trough and Portugal were sampled during annual-fisheries surveys run by Marine Science Scotland (MSS) and Instituto Português do Mar e da Atmosfera (IPMA) respectively. Additional samples were collected in Portugal from bycatches of the commercial black scabbardfish (Aphanopus carbo) longline and of the crustacean bottom trawl fisheries. In the Mediterranean Sea, E. spinax samples from the Balearic Islands were obtained from research cruises conducted by the Mediterranean International Trawl Survey (MEDITS). In the

114	Raunefjord (Norway), sharks were caught using deep-water longlines during scientific surveys.
115	Each collected specimen was sexed and measured for total length (TL). Following dissection,
116	white muscle was quickly frozen before analysis.
117	Samples were dried (48 hours at 60°C) and ground using pestle and mortar prior analysis. To
118	avoid possible biases linked to polar compound contents (i.e. lipids, urea and trimethylamine
119	$\emph{N}\text{-}\text{oxide}$), all samples were washed in distilled water and their $\delta^{13}\text{C}$ mathematically corrected
120	(Kiljunen et al., 2006; Li et al., 2016). Only samples from Norway obtained in 2017 underwent
121	lipid chemical extraction using a modified Folch method based on repetitive wash in a 2:1
122	dichloromethane:methanol mix (Folch et al., 1957). Carbon and nitrogen isotope
123	compositions were subsequently compared between individuals sampled in Norway in 2014
124	(i.e. corrected δ^{13} C values) and in 2017 (i.e. δ^{13} C values measured after extraction) at the same
125	locations. They were all kept in the dataset because no significant difference in mean isotopic
126	values was observed. As pure protein samples exhibit a C:N ratio around 3.0 for shark muscle,
127	a good lipid, urea and trimethylamine N-oxide extraction for all samples were considered as
128	those with a C:N ratio lower than 4.0 (Hussey et al., 2012) leading to no discard or additional
129	chemical extractions.
130	Isotopic ratios (δ) were expressed in per mille (∞) following: $\delta X = [(R_{sample}/R_{standard}) - 1] \times 1000$
131	where X is 13 C or 15 N and R is 13 C/ 12 C or 15 N/ 14 N, respectively. δ^{13} C values are expressed with
132	reference to the Vienna Pee Dee Belemnite and $\delta^{15} N$ values are expressed relative to
133	Atmospheric air. Depending on sites and sampling year, samples were analysed in different
134	facilities. Samples from Norway obtained in 2017 were processed in Liege (Oceanology,
135	ULiege, Belgium) using a continuous flow isotope ratio mass spectrometer (Isoprime 100,
136	Isoprime, United Kingdom) coupled to a C-N-S elemental analyser (MicroVario, Elementar,

Germany). The certified substances, provided by the International Atomic Energy Agency IAEA (Vienna) were IAEA-CH-6 (sucrose) for δ^{13} C and IAEA-N-1 (ammonium sulfate) for δ^{15} N. Cod (Gadus morhua) muscle was used as a natural replicate showing precision of ± 0.05 % for δ^{13} C and \pm 0.23 % for $\delta^{15}N$ (based on the standard deviation of the replicate measurements). Stable isotope ratios of samples obtained along the southern Iberian coast of Portugal in 2015 were measured using a Thermo Scientific Delta V Advantage IRMS via Conflo IV interface at Marinnova – Marine and Environmental Innovation, Technology and Services (Portugal). Samples from Rockall were analysed in two separate laboratories at the Scottish University Environmental Research Council (SUERC) and at the OEA Labs (Exeter, United Kingdom). The remaining samples from 2014 (i.e. Norway, Portugal and Balearic Islands) were processed in Elemtex (Gunnislake, United Kingdom). SUERC samples were run on a continuous flow Elementar vario PYRO cube elemental analyser coupled with a Thermo Scientific Delta V plus isotope ratio mass spectrometer (SUERC, NERC LSMSF, East Kilbride facility). OEA Labs and Elemtex samples were run on a Thermo EA 110 elemental analyser linked to a Europa Scientific 2020 isotope ratio mass spectrometer running in continuous flow mode.

2.3. Data analysis

137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

159

To assess isotopic baseline effects (i.e. change of primary producer isotopic composition) in each sampling site, isotope values were extracted from predictive models established for δ^{13} C (Magozzi et al., 2017) and δ^{15} N (Somes et al., 2010). Baseline values were extracted from the shark specific sampling locations (details of the specific latitudes and longitudes can be found in supplementary information). To overcome the baseline effect when comparing sharks between locations, the modelled baseline (i.e. phytoplankton) value was subtracted from shark stable isotope composition for each individual geographic

coordinate: $\Delta X = \delta X_{shark}$ - δX_{phyto} where X is ¹³ C or ¹⁵ N. Modelled δ^{15} N baseline values inside
the fjord were not available and values from the adjacent North Sea were subsequently used.
All statistical analyses were performed on the open source software R (R Core Team, 2020).
For each site, the SIBER package (Jackson et al., 2011) was used to estimate the size of the
isotopic niches (i.e. using $\Delta^{13}C$ and $\Delta^{15}N$) and their associated Layman metrics (Layman et al.,
2007). Layman metrics were first used to characterize trophic niche space using four metrics.
Isotopic ranges (Δ^{13} C rg and Δ^{15} N rg) described the distance between the most 13 C- and 15 N-
enriched and most depleted individuals respectively, the total area (TA), the size of the
isotopic niche based on convex hull area and the mean distance to the centroid (CD), the mean
distance of each individual to the $\Delta^{13}\text{C}/\Delta^{15}\text{N}$ centroid. The other two metrics reflected trophic
redundancy (i.e. the relative position of individuals from one another inside their isotopic
niche). Mean nearest neighbor distance (NND and its standard deviation SDNND) measured
the overall density of individuals clustering in a way that smaller NND and SDNND would
describe small differences between individual isotopic values (Layman et al., 2007). The SIBER
package was finally used to calculate the standard ellipse area (SEA) encompassing 40% of the
bulk $\delta^{13} \text{C}/\delta^{15} \text{N}$ data at each site and the isotopic overlaps between them as a proportion of
the non-overlapping area of two given ellipses.
For between-site comparisons, data was first checked for normality by Shapiro-Wilk tests and
homoscedasticity by Bartlett's tests. As both conditions were not met, Kruskal-Wallis tests
followed by Conover-Iman (C-I) post-hoc tests were applied. Comparisons between sexes
were carried out using Student's t-tests or its non-parametric analogue, the Wilcoxon test.
Pearson correlation tests were used to assess linear correlations between Δ^{13} C, Δ^{15} N and TL.
Finally, an analysis of covariance (ANCOVA) was performed to test for slope differences

between linear regressions. For all statistical analyses, the significance threshold was set at p<0.05.

3. RESULTS

3.1. Stomach content analysis

In the Raunefjord, a set of 255 stomachs was analysed (on different specimens than for stable isotope analysis), among which 80 contained remains of prey items. Individuals that had food remains in their stomach ranged from 14.3 to 52.0 cm (TL) and included 51 females and 29 males. Euphausiacea was the major prey items of *E. spinax* with occurrence of *Meganyctiphanes norvegica*. Other prey included other crustaceans (mostly decapods), teleost fishes and cephalopods (Table 1). Until they reached 36.0 cm, Euphausiacea was the only prey item identified in the guts of *E. spinax*. Beyond this size, a diet shift was observed with occurrence of teleost fishes (such as *Scomber scombrus* and *Maurolicus muelleri*) and, to a lesser extent, decapods and cephalopods, therefore decreasing the importance of Euphausiacea overall.

3.2. Stable isotopes

Muscle tissues from 147 *E. spinax* individuals were recovered from the four different stations (Table 2). As fishing methodologies differed between stations, capture depth significantly varied among sampling sites ($\chi^2_{146,3}$ = 98.2, p<0.001). Samples from Norway were caught on average at a depth of 243 meters, which was significantly shallower than the other sites where catch depth was around 600 meters. *E. spinax* TL varied from 11.0 to 57.4 cm and was significantly different between sampling sites ($\chi^2_{146,3}$ = 74.2, p<0.001). Individuals from

204	Norway and Rockall were significantly larger than in Portugal and in the Balearic Islands (C-I
205	test, p<0.001 for all pairs of comparison). Females were larger than males considering the
206	entire dataset (W = 1860, p<0.01), as observed in Portugal (W = 148, p<0.05) and in Norway
207	(t = -6.91, p<0.001). No significant difference in TL was found between sexes in the Balearic
208	Islands and Rockall Trough.
209	Modelled phytoplanktonic baselines significantly differed among sites ($\chi^2_{146,3}$ = 101.4, p<0.001
210	for $\delta^{13}C_{phyto}$; $\chi^2_{146,3}$ = 142.5, p<0.001 for $\delta^{15}N_{phyto}$). $\delta^{13}C_{phyto}$ was significantly lower in the
211	Balearic Islands than in Rockall and Portugal, while Norway displayed higher values than all
212	other sites (C-I test, p<0.001). Concerning $\delta^{15}N_{phyto}$, all locations significantly differed (C-I test,
213	p<0.001, for all pairs of comparison) with Portugal being the most ¹⁵ N-enriched, followed by
214	the Balearic Islands, Rockall Trough and southwestern Norway.
215	Shark bulk isotopic values (i.e. $\delta^{13}C$ and $\delta^{15}N$) differed among sampling sites (Figure 2) with
216	the only overlapping SEAs (Standard Ellipse Areas) reported between <i>E. spinax</i> sampled in the
217	Portugal Iberian system and Rockall Trough (39.89%). $\Delta^{15}N$ values differed between locations
218	($\chi^2_{146,3}$ = 118.3, p<0.001; Figure 3a). Norway individuals exhibited the highest $\Delta^{15}N$ and
219	individuals sampled in Portugal the lowest (C-I test, p<0.001 for all pairs of comparison).
220	Rockall $\Delta^{15}N$ value was also significantly higher than in Portugal (C-I test, p<0.01). $\Delta^{13}C$ also
221	varied significantly among sampling locations ($\chi^2_{146,3}$ = 89.4, p<0.001; Figure 3b) with
222	individuals from the Balearic Islands having the highest $\Delta^{13}\text{C}$ and individuals from Norway the
223	lowest (C-I test, p<0.001 for all pairs of comparison). E. spinax sampled in Rockall Trough and
224	Portugal had similar $\Delta^{13}C$ (C-I test, p>0.05). At each location, there was no depth-related or
225	sex differences in $\Delta^{13}\text{C}$ and $\Delta^{15}\text{N}$ except in the Norwegian fjord where both values were
226	significantly higher in males.

Individuals from Portugal had a singular profile encompassing the highest Δ^{13} C rg, TA and CD suggesting higher isotopic diversity (Table 3). In Norway and in the Balearic Islands, *E. spinax* presented the smallest TA, CD and the lowest Δ^{13} C and Δ^{15} N rg. While displaying intermediate TA and CD, *E. spinax* sampled in Rockall exhibited the highest Δ^{15} N rg.

For all sampling sites, significant linear relations were established between Δ^{15} N and TL (Figure 4a). Δ^{15} N increased with TL in three sites: Portugal, Rockall and the Balearic Islands. *E. spinax* sampled in Portugal and Rockall had equivalent slopes while specimens from the Balearic Islands showed a flatter one (ANCOVA, F=5.83 p<0.05 for Rockall and F=21.32 p<0.001 for Portugal). Specimens in Norway were the only ones with a significant decrease in Δ^{15} N values with TL. Δ^{13} C increased with TL in the Balearic Islands and Rockall with no differences in slopes (Figure 4b). At each location, linear regressions did not significantly change between sexes or

4. DISCUSSION

with depth.

4.1. Trophic ecology of E. spinax

4.1.1. Prey composition (Stomach content analysis)

In Norway, *E. spinax* stomach content composition inside the Raunefjord matched previous reports in the surrounding North and Norwegian Sea (Bergstad et al., 2003; Klimpel et al., 2003). The diet was dominated by *M. norvegica*, the most abundant prey in the ecosystem (Bergstad et al., 2003; Klimpel et al., 2003), with an ontogenetic diet switch toward the consumption of teleost fishes. The only difference was a later switch observed in the fjord, probably due to a sampling difference with the two previous studies as sampled individuals inside the fjord were larger.

250

251

252

253

254

255

256

257

258

259

260

261

262

263

264

265

266

267

268

269

270

271

At all locations sampled for stable isotope analysis, previously published studies have highlighted an ontogenetic shift in the diet of E. spinax, except for two studies on Portugal and the Balearic Islands (Table 4). Before the ontogenetic diet shift, Euphausiacea was generally the major prey item, mostly represented by the northern krill, Meganyctiphanes norvegica. Ontogenetic diet shift occurred at different lengths depending on the site, from 19.4 cm in Norway to 39.0 cm in Rockall Trough. Beyond these sizes, Euphausiacea decreased in importance due to the increasing occurrence of teleost fishes, cephalopods, or other crustaceans. This result was in line with previous analyses of E. spinax diet in the Atlantic Ocean and Mediterranean Sea (Bengil et al., 2019; Isbert et al., 2015; Preciado et al., 2017, 2009). In Rockall, the later shift toward the consumption of the decapod crustacean, Pasiphaea tarda, has been explained by larger individuals foraging closer to the sea bottom where prey availability is different (Mauchline and Gordon, 1983). Ontogenetic diet shifts are commonly observed in mesopredator fishes due to increasing body length, mouth gap and stomach size, hunting capacities and energetic demands, or to avoid intra-specific competition (Klimpel et al., 2003; Neiva et al., 2006). Stomach contents analysed in the Balearic Islands showed contrasting results between studies. E. spinax either foraged following the previously described ontogenetic shift from small Euphausiacea to teleost fishes (Fanelli et al., 2009; Macpherson, 1980) or fed on cephalopods throughout their entire size range in more recent studies (Valls et al., 2017, 2011). A higher proportion of cephalopod consumption by E. spinax was previously observed in the adjacent Catalan Sea and has been linked to the high exploitation rate of the area, depleting fish stocks in favour of cephalopods (Barría et al., 2018; Doubleday et al., 2016). In deep-sea sharks, smaller adult size and earlier maturity due to oligotrophic conditions in the

273

274

275

276

277

278

279

280

281

282

283

284

285

286

287

288

289

290

291

292

293

294

Mediterranean Sea could also explain cephalopod consumption by small specimens (Catarino et al., 2015; Massutí et al., 2004). However, this trend was not yet observed in *E. spinax* (Coelho et al., 2010).

4.1.2. Nitrogen and carbon isotope compositions ($\Delta^{15}N$ and $\Delta^{13}C$)

Stable isotope values obtained in this study were measured in multiple laboratories which may result in a potential bias (Mill et al., 2008). The analysis of a reference sample (USGS 40 glutamic acid) to ensure comparability was performed on two of the four instruments used in this study (i.e. for Elemtex and OEA laboratories). δ^{13} C values were on average 0.23 ± 0.40 % higher from Elemtex than those obtained from OEA Labs. This difference, although significant, is quite small in regard of isotope patterns interpreted in this study, and comparable to error estimates considered acceptable in many ecological studies using stable isotope tracers. Nevertheless, this difference was subtracted from all shark δ^{13} C values measured by Elemtex. There were no observed differences in $\delta^{15}N$ values. Specimens sampled in Norway in 2017 and in Portugal in 2015 were analysed in facilities where glutamic acid samples could not be measured. However, all facilities follow recommended procedures for stable isotope analyses, and use certified materials that are all calibrated against the same international references (Vienna Pee Dee Belemnite for δ^{13} C, Atmospheric air for δ^{15} N). Therefore, we argue that analytical biases are unlikely to have a major impact on stable isotope trends depicted here. Ecological differences in E. spinax isotopic niches between sites were analysed by standardising values (i.e. $\Delta^{15}N$ and $\Delta^{13}C$) to mitigate baseline effects (Bird et al., 2018). The absence of significant differences in isotopic ratios between females and males suggests a uniform diet across sexes, as globally observed in stomach content analyses of the species.

Nitrogen isotope composition is commonly used as a proxy of trophic position in marine food
web (Cabana and Rasmussen, 1994). M. norvegica, the main prey of E. spinax, is a low trophic
level omnivorous species feeding on both phytoplankton and copepods (Gomes et al., 2001;
Kaartvedt et al., 2002). It is therefore unlikely that divergences in food web length might have
contributed to the marked $\Delta^{15}N$ differences. Moreover, the suspected $\Delta^{15}N$ switch with TL was
not observed at all sites. The relevance of $\Delta^{\rm 15}N$ as a good proxy of the trophic level could be
conditioned by the occurrence of a clear ontogenetic change as observed in Portugal and
Rockall sharks. Along the Iberian slope, the rapid sedimentation of phytoplankton (Lopez-
Lopez et al., 2017) results in a direct linear energy flow, with phytoplankton mainly consumed
by Euphausiacea, themselves consumed by secondary predators such as juvenile <i>E. spinax</i>
(Gomes et al., 2001). Older E. spinax fed on higher trophic level species such as Pasiphaea
sivado or Micromesistius poutassou (Neiva et al., 2006; Santos and Borges, 2001) explaining
the $^{15}\text{N-enrichment}$ (Figure 4a). In Rockall the same phenomenon is suspected to occur, with
$\it M.~norvegica$ being replaced by possibly ^{15}N -enriched prey like the decapod crustacean $\it P.$
tarda or the teleost M. muelleri (Mauchline and Gordon, 1983). However, more recent
information on <i>E. spinax</i> diet are needed to confirm this effect as the only stomach content
description available came from samples obtained between 1973 and 1981 (Mauchline and
Gordon, 1983) and major environmental and anthropogenic changes could have modified the
shark diet over time.
In benthopelagic communities, the higher degradation of sinking particles is leading to higher
$\delta^{15} N$ values in deeper species (Trueman et al., 2014). Bathymetric segregation at the inter-
and intra-specific levels is commonly observed among deep-sea sharks (Clarke et al., 2005;
Neat et al., 2015) and has been reported for <i>E. spinax</i> (Coelho and Erzini, 2010). At each site,

changes in capture depth were not associated with an increase in $\delta^{15}N$ or $\Delta^{15}N$ values. 318 However, differences in E. spinax vertical segregation of maturity stages or bathymetric 319 constraint among sites could explain $\Delta^{15}N$ differences and would need to be specifically 320 addressed in the future. 321 Except for the Balearic Islands, E. spinax likely shared a pelagic habitat owing to their lower 322 323 Δ^{13} C values (Figure 3b). This is in accordance with the high rate of phytoplanktonic production fuelling the deeper layers of Portugal and Rockall continental slope systems (Gomes et al., 324 325 2001; Mauchline and Gordon, 1983). In the Raunefjord, E. spinax Δ^{13} C values agree with a diet 326 dominated by pelagic preys (Bergstad et al., 2003; Klimpel et al., 2003). E. spinax sampled in the Balearic Islands exhibited higher Δ^{13} C values (Figure 3b), suggesting 327 reliance on different forms of primary production, such as ¹³C-enriched benthic prey (Madurell 328 et al., 2008). It could correspond to the higher occurrence of cephalopods in E. spinax diet at 329 330 that site (i.e. mainly Teuthoidea and Sepioidea). These cephalopods exhibit an ontogenetic diet shift from benthic to pelagic prey (Valls et al., 2017, 2011). As scavenging was not reported 331 in the area and because mature cephalopods exceed E. spinax length, sharks are expected to 332

4.2. Habitat characteristics influence on isotope compositions

forage on juvenile early benthic life stages possibly explaining their Δ^{13} C values.

333

334

335

336

337

338

339

E. spinax in the Iberian slope ecosystems and Rockall Trough, even if mainly relying on pelagic production, presented similarly high indices of isotopic diversity (Table 3). Both sites are large continuous continental slopes (Mauchline and Gordon, 1991; Ribeiro et al., 2005) and topographic similarities might drive these similitudes. Indeed, ichthyofauna inhabiting continental slope ecosystems are known to integrate nutrient from a mixture of pelagic and

340 benthic origins (Mauchline and Gordon, 1991; Trueman et al., 2014). This trophic diversity is possibly further enhanced by the access to different topographic features (e.g. canyons, banks 341 342 or steep slopes) (Romero-Romero et al., 2016; Rowden et al., 2010). 343 In Portugal, from spring to late summer, changes in wind-driven mesoscale currents induce an upwelling of nutrient-rich cold water (Loureiro et al., 2005) that influences shelf and slope 344 ecosystems over great distances (Pérez et al., 2010; Ribeiro et al., 2005) and resulting in 345 346 shifting isotopic signals. Due to upwelling seasonality and geographical influence, temporal and spatial variations in δ^{13} C values are observed in species at the base of the food web (Lopez-347 348 Lopez et al., 2017). The small-scale variability in the activity of the upwelling associated with the spatial scale at which individuals were fished might contribute to the overall isotopic 349 variability, a dynamic also observed in Rockall where the pelagic production is seasonally 350 351 stimulated over the shelf area (White et al., 2005). 352 The Balearic Islands deep ecosystem is characterized by a smaller continental slope, deep escarpments and canyons (Acosta et al., 2003). While topographic conditions might be similar 353 to large continental slope systems, the main difference resides in the oligotrophic nature of 354 355 the Mediterranean Sea (Bosc et al., 2004; Estrada, 1996). Such conditions might result in the observed small isotopic niche due to limited $\Delta^{15}N$ variations and high redundancy index (Table 356 3). Still, their broad Δ^{13} C rg, with some 13 C-depleted specimens, might reveal the existence of 357 358 resource partitioning within the population, which could be fuelled by a diversity of prey of both benthic and pelagic origin (Albo-Puigserver et al., 2015; Madurell et al., 2008; Newman 359 et al., 2011). Such coupling between reservoirs and reliance on other habitat is probably a 360 response to oligotrophic conditions reducing pelagic production and prey availability (Valls et 361 al., 2014). Sharks could also forage inside canyons where higher benthic prey densities are 362

364

365

366

367

368

369

370

371

372

373

374

375

376

377

378

379

380

381

382

383

384

present (Massutí et al., 2004). Other ¹³C-enriched sources have been identified in the Balearic Islands such as food falls (Cartes et al., 2016) or seagrass exportation at deeper layers (Boudouresque et al., 2016) but this remains speculative and will need further studies to investigate. Water temperature, higher in the Mediterranean Sea compared to the other locations, could also have led to differences in isotopic values compared to the other locations. Indeed, temperature can affect isotopic turnover rates and discrimination factors (Bloomfield et al., 2011), beyond latitudinal baseline δ^{13} C and δ^{15} N variations (Magozzi et al., 2017). Fjords are narrow, deep ecosystems delimited by steep flanking slopes (Harris, 2012). Therefore, possible bathymetric constraints (Coelho and Erzini, 2010) might reduce E. spinax foraging habitat diversity and explain isotopic niche limited space and high redundancy. $\Delta^{15}N$ values (Figure 3a) may result from the important particle residency time (Saino and Hattori, 1980) in deep basins. Indeed, even if fjords are dynamic systems with quick surface water turnover (Asplin et al., 1999), they are stratified with sometimes limited exchanges between reservoirs (Aure et al., 1996). As sinking particles are trapped in deep layers, especially in fjords, they would exhibit higher baseline $\delta^{15}N$ values than suggested by surface model values (Saino and Hattori, 1980; Trueman et al., 2014). Even if experiencing an ontogenetic shift toward higher trophic position preys, E. spinax $\Delta^{15}N$ decreased with TL (Figure 4a). Sharks sampled in the Raunefjord were only mature individuals probably post-ontogenetic diet shift. This decrease could result from a shift in feeding habitat with early maturing sharks foraging strictly in fjords while larger sharks could forage outside in offshore areas. This hypothesis would ultimately lead to the observed decreasing $\Delta^{15}N$ values with length, as modelled $\delta^{15}N$ baseline was extracted from outside of the fjord. Another possibility is that large sharks

change their foraging behaviour towards active predation of smaller organisms with lower trophic positions, although this is not supported by the stomach content analysis.

Finally, relatively small deep-sea sharks can exhibit important horizontal migrations (Catarino et al., 2015; Rodríguez-Cabello and Sánchez, 2014). E. spinax population structure suggested it might connect distant areas across the northeastern Atlantic with a potential isolation of Mediterranean individuals (Gubili et al., 2016; McMillan et al., 2017). Migrating E. spinax are likely to feed on different isotopic baselines and would integrate them throughout the course of their migration (Carlisle et al., 2012). In this study, a significant overlap in bulk isotope niches occurred between Portugal and Rockall samples (Figure 2), reinforcing the idea of potential large-scale migration of E. spinax in the Atlantic Ocean. Conversely, the nonoverlapping and reduced isotopic niche spaces observed in the Balearic Islands and in the fjord suggest a certain level of residency, even more when diet is found homogeneous among distant sites. This strengthens the hypothesis of a separated population in the Mediterranean Sea, probably due to the bathymetric limitation at the Strait of Gibraltar (Catarino et al., 2015; Gubili et al., 2016). In Norway, while changes in $\Delta^{15}N$ values have been hypothetically linked to migrations outside the fjords, trophic redundancy and isotopic discrimination suggest E. spinax could be sedentary in the region. This hypothesis matches the separate stocks of E. spinax in Norwegian waters previously identified by vertebral chemistry variations (McMillan et al., 2017).

5. CONCLUSION

385

386

387

388

389

390

391

392

393

394

395

396

397

398

399

400

401

402

403

404

405

406

407

At each location, including inside the fjords, *Etmopterus spinax* appears as a benthopelagic mesopredator, mainly feeding on aggregations of *Meganyctiphanes norvegica* at a juvenile stage with increasing consumption of larger prey, such as teleost fishes, with

increasing length. Nonetheless, isotopic niche spaces varied significantly among sampling sites. Observed differences in carbon and nitrogen isotope compositions are likely to be explained by differences in habitat features (e.g. productive continental slopes vs oligotrophic conditions in the Mediterranean Sea) and other oceanographic characteristics (e.g. upwelling regime). Resulting variations in pelagic primary production rates and sequestration across the water column are likely to affect the strength of mesopelagic linkage toward deep reservoirs and subsequently *E. spinax* trophodynamic and isotopic values. Our results suggest that considering the influence of nutrient cycle on isotopic baselines allows a better understanding of the trophic ecology of predators in deep-sea habitats.

ACKNOWLEDGMENT

The authors would like to thank Christopher J. Somes, Sarah Magozzi and Katie St John Glew for providing the phytoplankton-modelled values, and to James Bell for his suggestions to early stages of the manuscript; as well as Tomas Sørlie, from the Espeland marine station, for the technical support during the shark sampling in Norway. This work was partially supported by FRIA grant (F.R.S-FNRS Belgium) to L.D. and N.P.. G.L and J.M are appointed by F.R.S-FNRS Belgium. The authors thank the Université de Bretagne Occidentale (UBO) and the École Doctorale des Sciences de la Mer et du Littoral (EDSML) for L.B. doctoral grant. This project is supported by ISblue project, interdisciplinary graduate school for the blue planet (ANR-17-EURE-0015) and co-funded by a grant from the French government under the program "Investissements d'Avenir". This study is the contribution BRC #374 of the Biodiversity Research Center (UCLouvain) from the Earth and Life Institute Biodiversity (ELIV) and the "Centre Interuniversitaire de Biologie Marine" (CIBIM). Thanks are also due to the

- 430 Portuguese Science Foundation for the financial support to R.P.V. (SFRH/BD/84030/2012) and
- CCMAR (UIDB/04326/2020) through national funds. 431

REFERENCES

432

447

448

449

450

451

452

453

454

455

456

457

458

459

460

461

- 433 Acosta, J., Canals, M., López-Martínez, J., Muñoz, A., Herranz, P., Urgeles, R., Palomo, C., Casamor, 434 J.L., 2003. The Balearic Promontory geomorphology (western Mediterranean): 435 morphostructure and active processes. Geomorphology 49, 177–204. 436 https://doi.org/10.1016/S0169-555X(02)00168-X
- 437 Albo-Puigserver, M., Navarro, J., Coll, M., Aguzzi, J., Cardona, L., Sáez-Liante, R., 2015. Feeding 438 ecology and trophic position of three sympatric demersal chondrichthyans in the 439 northwestern Mediterranean. Mar. Ecol. Prog. Ser. 524, 255–268. 440 https://doi.org/10.3354/meps11188
- Asplin, L., Salvanes, A.G.V., Kristoffersen, J.B., 1999. Nonlocal wind-driven fjord-coast advection and 441 its potential effect on plankton and fish recruitment. Fish. Oceanogr. 8, 255–263. 442 443 https://doi.org/10.1046/j.1365-2419.1999.00109.x
- 444 Aure, J., Molvær, J., Stigebrandt, A., 1996. Observations of inshore water exchange forced by a 445 fluctuating offshore density field. Mar. Pollut. Bull. 33, 112–119. 446 https://doi.org/10.1016/S0025-326X(97)00005-2
 - Bailey, D. m., Collins, M. a., Gordon, J. d. m., Zuur, A. f., Priede, I. g., 2009. Long-term changes in deep-water fish populations in the northeast Atlantic: a deeper reaching effect of fisheries? Proc. R. Soc. B Biol. Sci. 276, 1965–1969. https://doi.org/10.1098/rspb.2009.0098
 - Barría, C., Navarro, J., Coll, M., 2018. Feeding habits of four sympatric sharks in two deep-water fishery areas of the western Mediterranean Sea. Deep Sea Res. Part Oceanogr. Res. Pap. 142, 34-43. https://doi.org/10.1016/j.dsr.2018.09.010
 - Bengil, F., Bengil, E.G.T., Mavruk, S., Heral, O., Karaman, O.D., Ozaydin, O., 2019. Feeding Ecology of Four Demersal Shark Species (Etmopterus spinax, Galeus melastomus, Scyliorhinus canicula and Squalus blainville) from the Eastern Aegean Sea. Turk. J. Fish. Aquat. Sci. 19, 475–484.
 - Benn, A.R., Weaver, P.P., Billet, D.S.M., Hove, S. van den, Murdock, A.P., Doneghan, G.B., Bas, T.L., 2010. Human Activities on the Deep Seafloor in the North East Atlantic: An Assessment of Spatial Extent. PLOS ONE 5, e12730. https://doi.org/10.1371/journal.pone.0012730
 - Bergstad, O., Wik, D., Hildre, O., 2003. Predator-Prey Relationships and Food Sources of the Skagerrak Deep-water Fish Assemblage. J. Northwest Atl. Fish. Sci. 31, 165–180. https://doi.org/10.2960/J.V31.A12
- 463 Bird, C.S., Veríssimo, A., Magozzi, S., Abrantes, K.G., Aguilar, A., Al-Reasi, H., Barnett, A., Bethea, 464 D.M., Biais, G., Borrell, A., Bouchoucha, M., Boyle, M., Brooks, E.J., Brunnschweiler, J., 465 Bustamante, P., Carlisle, A., Catarino, D., Caut, S., Cherel, Y., Chouvelon, T., Churchill, D., 466 Ciancio, J., Claes, J., Colaço, A., Courtney, D.L., Cresson, P., Daly, R., de Necker, L., Endo, T., 467 Figueiredo, I., Frisch, A.J., Hansen, J.H., Heithaus, M., Hussey, N.E., litembu, J., Juanes, F., 468 Kinney, M.J., Kiszka, J.J., Klarian, S.A., Kopp, D., Leaf, R., Li, Y., Lorrain, A., Madigan, D.J., 469 Maljković, A., Malpica-Cruz, L., Matich, P., Meekan, M.G., Ménard, F., Menezes, G.M., 470 Munroe, S.E.M., Newman, M.C., Papastamatiou, Y.P., Pethybridge, H., Plumlee, J.D., Polo-471 Silva, C., Quaeck-Davies, K., Raoult, V., Reum, J., Torres-Rojas, Y.E., Shiffman, D.S., Shipley, 472 O.N., Speed, C.W., Staudinger, M.D., Teffer, A.K., Tilley, A., Valls, M., Vaudo, J.J., Wai, T.-C.,
- 473 Wells, R.J.D., Wyatt, A.S.J., Yool, A., Trueman, C.N., 2018. A global perspective on the trophic

- Bloomfield, A.L., Elsdon, T.S., Walther, B.D., Gier, E.J., Gillanders, B.M., 2011. Temperature and diet affect carbon and nitrogen isotopes of fish muscle: can amino acid nitrogen isotopes explain effects? J. Exp. Mar. Biol. Ecol. 399, 48–59. https://doi.org/10.1016/j.jembe.2011.01.015
- Bosc, E., Bricaud, A., Antoine, D., 2004. Seasonal and interannual variability in algal biomass and
 primary production in the Mediterranean Sea, as derived from 4 years of SeaWiFS
 observations. Glob. Biogeochem. Cycles 18. https://doi.org/10.1029/2003GB002034

- Boudouresque, C.F., Pergent, G., Pergent-Martini, C., Ruitton, S., Thibaut, T., Verlaque, M., 2016. The necromass of the Posidonia oceanica seagrass meadow: fate, role, ecosystem services and vulnerability. Hydrobiologia 781, 25–42. https://doi.org/10.1007/s10750-015-2333-y
- Cabana, G., Rasmussen, J.B., 1994. Modelling food chain structure and contaminant bioaccumulation using stable nitrogen isotopes. Nature 372, 255–257. https://doi.org/10.1038/372255a0
- Carlisle, A.B., Kim, S.L., Semmens, B.X., Madigan, D.J., Jorgensen, S.J., Perle, C.R., Anderson, S.D., Chapple, T.K., Kanive, P.E., Block, B.A., 2012. Using Stable Isotope Analysis to Understand the Migration and Trophic Ecology of Northeastern Pacific White Sharks (Carcharodon carcharias). PLOS ONE 7, e30492. https://doi.org/10.1371/journal.pone.0030492
- Cartes, J.E., Soler-Membrives, A., Stefanescu, C., Lombarte, A., Carrassón, M., 2016. Contributions of allochthonous inputs of food to the diets of benthopelagic fish over the northwest Mediterranean slope (to 2300m). Deep Sea Res. Part Oceanogr. Res. Pap. 109, 123–136. https://doi.org/10.1016/j.dsr.2015.11.001
- Catarino, D., Knutsen, H., Veríssimo, A., Olsen, E.M., Jorde, P.E., Menezes, G., Sannæs, H., Stanković, D., Company, J.B., Neat, F., Danovaro, R., Dell'Anno, A., Rochowski, B., Stefanni, S., 2015. The Pillars of Hercules as a bathymetric barrier to gene flow promoting isolation in a global deepsea shark (Centroscymnus coelolepis). Mol. Ecol. 24, 6061–6079. https://doi.org/10.1111/mec.13453
- Chavez, F.P., Ryan, J., Lluch-Cota, S.E., C, M.Ñ., 2003. From Anchovies to Sardines and Back: Multidecadal Change in the Pacific Ocean. Science 299, 217–221. https://doi.org/10.1126/science.1075880
- Chouvelon, T., Spitz, J., Caurant, F., Mèndez-Fernandez, P., Chappuis, A., Laugier, F., Le Goff, E., Bustamante, P., 2012. Revisiting the use of δ 15N in meso-scale studies of marine food webs by considering spatio-temporal variations in stable isotopic signatures The case of an open ecosystem: The Bay of Biscay (North-East Atlantic). Prog. Oceanogr. 101, 92–105. https://doi.org/10.1016/j.pocean.2012.01.004
- Churchill, D.A., Heithaus, M.R., Vaudo, J.J., Grubbs, R.D., Gastrich, K., Castro, J.I., 2015. Trophic interactions of common elasmobranchs in deep-sea communities of the Gulf of Mexico revealed through stable isotope and stomach content analysis. Deep Sea Res. Part II Top. Stud. Oceanogr., Biology of Deep-Water Chondrichthyans 115, 92–102. https://doi.org/10.1016/j.dsr2.2014.10.011
- Claes, J.M., Aksnes, D.L., Mallefet, J., 2010. Phantom hunter of the fjords: Camouflage by counterillumination in a shark (Etmopterus spinax). J. Exp. Mar. Biol. Ecol. 388, 28–32. https://doi.org/10.1016/j.jembe.2010.03.009
- Clarke, M.W., Borges, L., Officer, R.A., 2005. Comparisons of Trawl and Longline Catches of Deepwater Elasmobranchs West and North of Ireland. J. Northwest Atl. Fish. Sci. 35, 429–442.
- Coelho, R., Erzini, K., 2010. Depth distribution of the velvet belly, Etmopterus spinax, in relation to
 growth and reproductive cycle: The case study of a deep-water lantern shark with a wide ranging critical habitat. Mar. Biol. Res. 6, 381–389.
 https://doi.org/10.1080/17451000802644706
- Coelho, R., Rey, J., Sola, L.G. de, Carvalho, J.F. de, Erzini, K., 2010. Comparing Atlantic and
 Mediterranean populations of the velvet belly lanternshark, Etmopterus spinax, with
 comments on the efficiency of density-dependent compensatory mechanisms. Mar. Biol. Res.
 6, 373–380. https://doi.org/10.1080/17451000903300885

- Cortés, E., 1999. Standardized diet compositions and trophic levels of sharks. ICES J. Mar. Sci. 56,
 707–717. https://doi.org/10.1006/jmsc.1999.0489
- Cortés, E., 1997. A critical review of methods of studying fish feeding based on analysis of stomach
 contents: application to elasmobranch fishes. Can. J. Fish. Aquat. Sci. 54, 726–738.
 https://doi.org/10.1139/f96-316
- Devine, J.A., Baker, K.D., Haedrich, R.L., 2006. Deep-sea fishes qualify as endangered. Nature 439, 29–29. https://doi.org/10.1038/439029a
- Doubleday, Z.A., Prowse, T.A.A., Arkhipkin, A., Pierce, G.J., Semmens, J., Steer, M., Leporati, S.C.,
 Lourenço, S., Quetglas, A., Sauer, W., Gillanders, B.M., 2016. Global proliferation of
 cephalopods. Curr. Biol. 26, R406–R407. https://doi.org/10.1016/j.cub.2016.04.002
- Duchatelet, L., Claes, J.M., Delroisse, J., Flammang, P., Mallefet, J., 2021. Glow on Sharks: State of the
 Art on Bioluminescence Research. Oceans 2, 822–842.
 https://doi.org/10.3390/oceans2040047
- 539 Estrada, M., 1996. Primary production in the northwestern Mediterranean. Sci. Mar. 60.

541

542

543

544

545

546

547

548

549

550

551

552

553

554

555

556

557558

- Fanelli, E., Rey, J., Torres, P., Sola, L.G. de, 2009. Feeding habits of blackmouth catshark Galeus melastomus Rafinesque, 1810 and velvet belly lantern shark Etmopterus spinax (Linnaeus, 1758) in the western Mediterranean. J. Appl. Ichthyol. 25, 83–93. https://doi.org/10.1111/j.1439-0426.2008.01112.x
- Folch, J., Lees, M., Sloane Stanley, G.H., 1957. A simple method for the isolation and purification of total lipides from animal tissues. J. Biol. Chem. 226, 497–509.
- Fry, B., Sherr, E.B., 1984. δ13C Measurements as Indicators of Carbon Flow in Marine and Freshwater Ecosystems, in: Rundel, P.W., Ehleringer, J.R., Nagy, K.A. (Eds.), Stable Isotopes in Ecological Research, Ecological Studies. Springer, New York, NY, pp. 196–229. https://doi.org/10.1007/978-1-4612-3498-2_12
- Gomes, M.C., Serrão, E., de Fátima Borges, M., 2001. Spatial patterns of groundfish assemblages on the continental shelf of Portugal. ICES J. Mar. Sci. 58, 633–647. https://doi.org/10.1006/jmsc.2001.1052
- Gubili, C., Macleod, K., Perry, W., Hanel, P., Batzakas, I., Farrell, E.D., Lynghammar, A., Mancusi, C., Mariani, S., Menezes, G.M., Neat, F., Scarcella, G., Griffiths, A.M., 2016. Connectivity in the deep: Phylogeography of the velvet belly lanternshark. Deep Sea Res. Part Oceanogr. Res. Pap. 115, 233–239. https://doi.org/10.1016/j.dsr.2016.07.002
- Harris, P.T., 2012. Seafloor Geomorphology—Coast, Shelf, and Abyss, in: Harris, P.T., Baker, E.K. (Eds.), Seafloor Geomorphology as Benthic Habitat. Elsevier, London, pp. 109–155. https://doi.org/10.1016/B978-0-12-385140-6.00006-2
- Heithaus, M.R., Frid, A., Wirsing, A.J., Worm, B., 2008. Predicting ecological consequences of marine
 top predator declines. Trends Ecol. Evol. 23, 202–210.
 https://doi.org/10.1016/j.tree.2008.01.003
- 563 Howell, K.L., Hilário, A., Allcock, A.L., Bailey, D., Baker, M., Clark, M.R., Colaço, A., Copley, J., Cordes, 564 E.E., Danovaro, R., Dissanayake, A., Escobar, E., Esquete, P., Gallagher, A.J., Gates, A.R., Gaudron, S.M., German, C.R., Gjerde, K.M., Higgs, N.D., Le Bris, N., Levin, L.A., Manea, E., 565 566 McClain, C., Menot, L., Mestre, N.C., Metaxas, A., Milligan, R., Muthumbi, A.W.N., 567 Narayanaswamy, B.E., Ramalho, S.P., Ramirez-Llodra, E., Robson, L.M., Rogers, A.D., Sellanes, 568 J., Sigwart, J.D., Sink, K., Snelgrove, P.V.R., Stefanoudis, P.V., Sumida, P.Y., Taylor, M.L., 569 Thurber, A.R., Vieira, R., Watanabe, H.K., Woodall, L.C., Xavier, J.R., 2021. A decade to study 570 deep-sea life. Nat. Ecol. Evol. 5, 265-267. https://doi.org/10.1038/s41559-020-01352-5
- Hussey, N.E., MacNeil, M.A., Olin, J.A., McMeans, B.C., Kinney, M.J., Chapman, D.D., Fisk, A.T., 2012.
 Stable isotopes and elasmobranchs: tissue types, methods, applications and assumptions. J.
 Fish Biol. 80, 1449–1484. https://doi.org/10.1111/j.1095-8649.2012.03251.x
- Hussey, N.E., Orr, J., Fisk, A.T., Hedges, K.J., Ferguson, S.H., Barkley, A.N., 2018. Mark report satellite tags (mrPATs) to detail large-scale horizontal movements of deep water species: First results

- for the Greenland shark (Somniosus microcephalus). Deep Sea Res. Part Oceanogr. Res. Pap. 134, 32–40. https://doi.org/10.1016/j.dsr.2018.03.002
- Isbert, W., Rodríguez-Cabello, C., Frutos, I., Preciado, I., Montero, F.E., Pérez-del-Olmo, A., 2015.
 Metazoan parasite communities and diet of the velvet belly lantern shark Etmopterus spinax
 (Squaliformes: Etmopteridae): a comparison of two deep-sea ecosystems. J. Fish Biol. 86,
 687–706. https://doi.org/10.1111/jfb.12591

- Jackson, A.L., Inger, R., Parnell, A.C., Bearhop, S., 2011. Comparing isotopic niche widths among and within communities: SIBER Stable Isotope Bayesian Ellipses in R. J. Anim. Ecol. 80, 595–602. https://doi.org/10.1111/j.1365-2656.2011.01806.x
- Kaartvedt, S., Larsen, T., Hjelmseth, K., Onsrud, M.S.R., 2002. Is the omnivorous krill Meganyctiphanes norvegica primarily a selectively feeding carnivore? Mar. Ecol. Prog. Ser. 228, 193–204. https://doi.org/10.3354/meps228193
- Kiljunen, M., Grey, J., Sinisalo, T., Harrod, C., Immonen, H., Jones, R.I., 2006. A revised model for lipid-normalizing δ13C values from aquatic organisms, with implications for isotope mixing models. J. Appl. Ecol. 43, 1213–1222. https://doi.org/10.1111/j.1365-2664.2006.01224.x
- Klimpel, S., Palm, H.W., Seehagen, A., 2003. Metazoan parasites and food composition of juvenile Etmopterus spinax (L., 1758) (Dalatiidae, Squaliformes) from the Norwegian Deep. Parasitol. Res. 89, 245–251. https://doi.org/10.1007/s00436-002-0741-1
- Layman, C.A., Araujo, M.S., Boucek, R., Hammerschlag-Peyer, C.M., Harrison, E., Jud, Z.R., Matich, P., Rosenblatt, A.E., Vaudo, J.J., Yeager, L.A., Post, D.M., Bearhop, S., 2012. Applying stable isotopes to examine food-web structure: an overview of analytical tools. Biol. Rev. 87, 545–562. https://doi.org/10.1111/j.1469-185X.2011.00208.x
- Layman, C.A., Arrington, D.A., Montaña, C.G., Post, D.M., 2007. Can Stable Isotope Ratios Provide for Community-Wide Measures of Trophic Structure? Ecology 88, 42–48. https://doi.org/10.1890/0012-9658(2007)88[42:CSIRPF]2.0.CO;2
- Li, Y., Zhang, Y., Hussey, N.E., Dai, X., 2016. Urea and lipid extraction treatment effects on δ15N and δ13C values in pelagic sharks. Rapid Commun. Mass Spectrom. 30, 1–8. https://doi.org/10.1002/rcm.7396
- Lopez-Lopez, L., Preciado, I., Muñoz, I., Decima, M., Molinero, J.C., Tel, E., 2017. Does upwelling intensity influence feeding habits and trophic position of planktivorous fish? Deep Sea Res. Part Oceanogr. Res. Pap. 122, 29–40. https://doi.org/10.1016/j.dsr.2017.01.017
- Lorrain, A., Graham, B.S., Popp, B.N., Allain, V., Olson, R.J., Hunt, B.P.V., Potier, M., Fry, B., Galván-Magaña, F., Menkes, C.E.R., Kaehler, S., Ménard, F., 2015. Nitrogen isotopic baselines and implications for estimating foraging habitat and trophic position of yellowfin tuna in the Indian and Pacific Oceans. Deep Sea Res. Part II Top. Stud. Oceanogr., Impacts of climate on marine top predators 113, 188–198. https://doi.org/10.1016/j.dsr2.2014.02.003
- Loureiro, S., Newton, A., Icely, J., 2005. Microplankton composition, production and upwelling dynamics in Sagres (SW Portugal) during summer of 2001. Sientia Mar. 69, 323–341.
- Macpherson, E., 1980. Régime alimentaire de Galeus melastomus (Rafinesque, 1810), Etmopterus 8 spinax (L., 1758) et Scymnorhinus licha (Bonaterre, 1788), en Méditerranée Occidentale. Vie Milieu 30, 139–148.
- Madurell, T., Fanelli, E., Cartes, J.E., 2008. Isotopic composition of carbon and nitrogen of
 suprabenthic fauna in the NW Balearic Islands (western Mediterranean). J. Mar. Syst., The
 Wrapping Up of the IDEA Project: 71, 336–345.
 https://doi.org/10.1016/j.jmarsys.2007.03.006
- Magozzi, S., Yool, A., Zanden, H.B.V., Wunder, M.B., Trueman, C.N., 2017. Using ocean models to predict spatial and temporal variation in marine carbon isotopes. Ecosphere 8, e01763. https://doi.org/10.1002/ecs2.1763
- Massutí, E., Gordon, J., Moranta, J., Swan, S., Stefanescu, C., Merrett, N.R., 2004. Mediterranean and Atlantic deep-sea fish assemblages: Differences in biomass composition and size-related structure. Sci. Mar. 68, 101–115.

- Mauchline, J., Gordon, J., 1991. Oceanic prey of benthopelagic fish in the benthic boundary layer of a marginal oceanic region. Mar. Ecol.-Prog. Ser. - MAR ECOL-PROGR SER 74, 109–115. https://doi.org/10.3354/meps074109
- Mauchline, J., Gordon, J.D.M., 1983. Diets of the sharks and chimaeroids of the Rockall Trough, northeastern Atlantic Ocean. Mar. Biol. 75, 269–278. https://doi.org/10.1007/BF00406012
- 632 McMillan, M.N., Izzo, C., Junge, C., Albert, O.T., Jung, A., Gillanders, B.M., 2017. Analysis of vertebral 633 chemistry to assess stock structure in a deep-sea shark, Etmopterus spinax. ICES J. Mar. Sci. 634 74, 793–803. https://doi.org/10.1093/icesjms/fsw176
- Mill, A.C., Sweeting, C.J., Barnes, C., Al-Habsi, S.H., MacNeil, M.A., 2008. Mass-spectrometer bias in
 stable isotope ecology. Limnol. Oceanogr. Methods 6, 34–39.
 https://doi.org/10.4319/lom.2008.6.34
- 638 Moura, T., Fernandes, A., Figueiredo, I., Alpoim, R., Azevedo, M., 2018. Management of deep-water 639 sharks' by-catch in the Portuguese anglerfish fishery: from EU regulations to practice. Mar. 640 Policy 90, 55–67. https://doi.org/10.1016/j.marpol.2018.01.006

642

643

644

645

646

647

648

649

650

651

652

653

654

655

656 657

658

659

660

661

662

663

664 665

666 667

668

669 670

- Neat, F.C., Burns, F., Jones, E., Blasdale, T., 2015. The diversity, distribution and status of deep-water elasmobranchs in the Rockall Trough, north-east Atlantic Ocean. J. Fish Biol. 87, 1469–1488. https://doi.org/10.1111/jfb.12822
- Neiva, J., Coelho, R., Erzini, K., 2006. Feeding habits of the velvet belly lanternshark Etmopterus spinax (Chondrichthyes: Etmopteridae) off the Algarve, southern Portugal. J. Mar. Biol. Assoc. U. K. 86, 835–841. https://doi.org/10.1017/S0025315406013762
- Newman, M.C., Xu, X., Cotton, C.F., Tom, K.R., 2011. High Mercury Concentrations Reflect Trophic Ecology of Three Deep-Water Chondrichthyans. Arch. Environ. Contam. Toxicol. 60, 618–625. https://doi.org/10.1007/s00244-010-9584-4
- Newsome, S.D., Rio, C.M. del, Bearhop, S., Phillips, D.L., 2007. A Niche for Isotopic Ecology. Front. Ecol. Environ. 5, 429–436.
- Pérez, F.F., Padín, X.A., Pazos, Y., Gilcoto, M., Cabanas, M., Pardo, P.C., Doval, M.D., Farina-Busto, L., 2010. Plankton response to weakening of the Iberian coastal upwelling. Glob. Change Biol. 16, 1258–1267. https://doi.org/10.1111/j.1365-2486.2009.02125.x
- Pinsky, M.L., Jensen, O.P., Ricard, D., Palumbi, S.R., 2011. Unexpected patterns of fisheries collapse in the world's oceans. Proc. Natl. Acad. Sci. 108, 8317–8322. https://doi.org/10.1073/pnas.1015313108
- Polunin, N., Morales-Nin, B., Pawsey, W.E., Cartes, J., Pinnegar, J., Moranta, J., 2001. Feeding relationships in Mediterranean bathyal assemblages elucidated by nitrogen and carbon isotope data. Mar. Ecol. Prog. Ser. 220, 13–23. https://doi.org/10.3354/meps220013
- Post, D.M., 2002. Using Stable Isotopes to Estimate Trophic Position: Models, Methods, and Assumptions. Ecology 83, 703–718. https://doi.org/10.1890/0012-9658(2002)083[0703:USITET]2.0.CO;2
- Preciado, I., Cartes, J.E., Punzón, A., Frutos, I., López-López, L., Serrano, A., 2017. Food web functioning of the benthopelagic community in a deep-sea seamount based on diet and stable isotope analyses. Deep Sea Res. Part II Top. Stud. Oceanogr., Advances in deep-sea biology: biodiversity, ecosystem functioning and conservation 137, 56–68. https://doi.org/10.1016/j.dsr2.2016.07.013
- Preciado, I., Cartes, J.E., Serrano, A., Velasco, F., Olaso, I., Sánchez, F., Frutos, I., 2009. Resource utilization by deep-sea sharks at the Le Danois Bank, Cantabrian Sea, north-east Atlantic Ocean. J. Fish Biol. 75, 1331–1355. https://doi.org/10.1111/j.1095-8649.2009.02367.x
- 672 Priede, I., Godbold, J., Niedzielski, T., Collins, M., Bailey, D., Gordon, J., Zuur, A., 2011. A review of the 673 spatial extent of fishery effects and species vulnerability of the deep-sea demersal fish 674 assemblage of the Porcupine Seabight, Northeast Atlantic Ocean (ICES Subarea VII). ICES J.
- 675 Mar. Sci. 68, 281–289. https://doi.org/10.1093/icesjms/fsq045

- Puccinelli, E., McQuaid, C.D., Dobretsov, S., Christofoletti, R.A., 2019. Coastal upwelling affects filter feeder stable isotope composition across three continents. Mar. Environ. Res. 147, 13–23.
 https://doi.org/10.1016/j.marenvres.2019.03.015
- R Core Team, 2020. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.

- Rau, G.H., Riebesell, U., Wolf-Gladrow, D., 1997. CO2aq-dependent photosynthetic 13C fractionation in the ocean: A model versus measurements. Glob. Biogeochem. Cycles 11, 267–278. https://doi.org/10.1029/97GB00328
 - Ribeiro, A.C., Peliz, Á., Santos, A.M.P., 2005. A study of the response of chlorophyll-a biomass to a winter upwelling event off Western Iberia using SeaWiFS and in situ data. J. Mar. Syst. 53, 87–107. https://doi.org/10.1016/j.jmarsys.2004.05.031
 - Rodríguez-Cabello, C., Sánchez, F., 2014. Is Centrophorus squamosus a highly migratory deep-water shark? Deep Sea Res. Part Oceanogr. Res. Pap. 92, 1–10. https://doi.org/10.1016/j.dsr.2014.06.005
 - Romero-Romero, S., Molina-Ramírez, A., Höfer, J., Duineveld, G., Rumín-Caparrós, A., Sanchez-Vidal, A., Canals, M., Acuña, J.L., 2016. Seasonal pathways of organic matter within the Avilés submarine canyon: Food web implications. Deep Sea Res. Part Oceanogr. Res. Pap. 117, 1–10. https://doi.org/10.1016/j.dsr.2016.09.003
 - Rowden, A.A., Schlacher, T.A., Williams, A., Clark, M.R., Stewart, R., Althaus, F., Bowden, D.A., Consalvey, M., Robinson, W., Dowdney, J., 2010. A test of the seamount oasis hypothesis: seamounts support higher epibenthic megafaunal biomass than adjacent slopes. Mar. Ecol. 31, 95–106. https://doi.org/10.1111/j.1439-0485.2010.00369.x
 - Saino, T., Hattori, A., 1980. 15 N natural abundance in oceanic suspended particulate matter. Nature 283, 752–754. https://doi.org/10.1038/283752a0
 - Santos, J., Borges, T., 2001. Trophic relationships in deep-water fish communities off Algarve, Portugal. Fish. Res., DEEP-WATER FISH AND FISHERIES: A SELECTION OF PAPERS PRESENTED AT THE ICES ANNUAL SCIENCE CONFERENCE IN LISBON, PORTUGAL 16-18 SEPTEMBER 1998. 51, 337–341. https://doi.org/10.1016/S0165-7836(01)00257-0
 - Shipley, Oliver.N., Brooks, E.J., Madigan, D.J., Sweeting, C.J., Dean Grubbs, R., 2017. Stable isotope analysis in deep-sea chondrichthyans: recent challenges, ecological insights, and future directions. Rev. Fish Biol. Fish. 27, 481–497. https://doi.org/10.1007/s11160-017-9466-1
 - Shipley, O.N., Polunin, N.V.C., Newman, S.P., Sweeting, C.J., Barker, S., Witt, M.J., Brooks, E.J., 2017. Stable isotopes reveal food web dynamics of a data-poor deep-sea island slope community. Food Webs 10, 22–25. https://doi.org/10.1016/j.fooweb.2017.02.004
 - Simpfendorfer, C.A., Kyne, P.M., 2009. Limited potential to recover from overfishing raises concerns for deep-sea sharks, rays and chimaeras. Environ. Conserv. 36, 97–103. https://doi.org/10.1017/S0376892909990191
 - Somes, C.J., Schmittner, A., Galbraith, E.D., Lehmann, M.F., Altabet, M.A., Montoya, J.P., Letelier, R.M., Mix, A.C., Bourbonnais, A., Eby, M., 2010. Simulating the global distribution of nitrogen isotopes in the ocean. Glob. Biogeochem. Cycles 24. https://doi.org/10.1029/2009GB003767
- 716 Thurber, A.R., Sweetman, A.K., Narayanaswamy, B.E., Jones, D.O.B., Ingels, J., Hansman, R.L., 2014. 717 Ecosystem function and services provided by the deep sea. Biogeosciences 11, 3941–3963. 718 https://doi.org/10.5194/bg-11-3941-2014
- Trueman, C.N., Johnston, G., O'Hea, B., MacKenzie, K.M., 2014. Trophic interactions of fish
 communities at midwater depths enhance long-term carbon storage and benthic production
 on continental slopes. Proc. R. Soc. B Biol. Sci. 281, 20140669.
 https://doi.org/10.1098/rspb.2014.0669
- Tu, C.-Y., Chen, K.-T., Hsieh, C., 2018. Fishing and temperature effects on the size structure of exploited fish stocks. Sci. Rep. 8, 1–10. https://doi.org/10.1038/s41598-018-25403-x

725	Valls, M., Quetglas, A., Moranta, J., Ordines, F., 2011. Feeding ecology of demersal elasmobranchs
726	from the shelf and slope off the Balearic Sea (western Mediterranean). Sci. Mar. 75, 633-
727	639. https://doi.org/10.3989/scimar.2011.75n4633
728	Valls, M., Rueda, L., Quetglas, A., 2017. Feeding strategies and resource partitioning among
729	elasmobranchs and cephalopods in Mediterranean deep-sea ecosystems. Deep Sea Res. Part
730	Oceanogr. Res. Pap. 128, 28-41. https://doi.org/10.1016/j.dsr.2017.09.002
731	Valls, M., Sweeting, C.J., Olivar, M.P., Fernández de Puelles, M.L., Pasqual, C., Polunin, N.V.C.,
732	Quetglas, A., 2014. Structure and dynamics of food webs in the water column on shelf and
733	slope grounds of the western Mediterranean. J. Mar. Syst., The wrapping up of the IDEADOS
734	project: International Workshop on Environment, Ecosystems and Demersal Resources, and
735	Fisheries 138, 171–181. https://doi.org/10.1016/j.jmarsys.2014.04.002
736	Vieira, R.P., Bett, B.J., Jones, D.O.B., Durden, J.M., Morris, K.J., Cunha, M.R., Trueman, C.N., Ruhl,
737	H.A., 2020. Deep-sea sponge aggregations (Pheronema carpenteri) in the Porcupine Seabigh
738	(NE Atlantic) potentially degraded by demersal fishing. Prog. Oceanogr. 183, 102189.
739	https://doi.org/10.1016/j.pocean.2019.102189
740	Vieira, R.P., Trueman, C.N., Readdy, L., Kenny, A., Pinnegar, J.K., 2019. Deep-water fisheries along the
741	British Isles continental slopes: status, ecosystem effects and future perspectives. J. Fish Biol
742	94, 981–992. https://doi.org/10.1111/jfb.13927
743	White, M., Mohn, C., de Stigter, H., Mottram, G., 2005. Deep-water coral development as a function
744	of hydrodynamics and surface productivity around the submarine banks of the Rockall
745	Trough, NE Atlantic, in: Freiwald, A., Roberts, J.M. (Eds.), Cold-Water Corals and Ecosystems,
746	Erlangen Earth Conference Series. Springer, Berlin, Heidelberg, pp. 503–514.
747	https://doi.org/10.1007/3-540-27673-4_25
748	

Journal Pre-proof

<u>Table 1:</u> Number (N), numeric index (%N), gravimetric index (%W), occurrence index (%O) and index of relative importance (%IRI) for each prey item found in stomachs of *Etmopterus spinax* sampled from 2008 to 2012 in the Raunefjord (Norway).

Prey items	N	%N	%W	%O	%IRI
Euphausiacea	39	41.94	7.04	55.10	45.55
Meganyctiphanes norvegica	12	12.90	3.29	18.37	7.63
Euphausiacea unidentified	27	29.03	3.75	44.90	37.75
Decapoda	7	7.53	17.17	14.29	5.95
Pasiphaea sivado	2	2.15	1.71	4.08	0.40
Pasiphaea multidentata	1	1.08	1.28	2.04	0.12
Pandalus montagui	1	1.08	11.78	2.04	0.67
Decapoda unidentified	3	3.23	2.40	6.12	0.88
Crustacea unidentified	10	10.75	5.92	20.41	5.74
Teleostei	7	7.53	52.76	14.29	14.54
Scomber scombrus	1	1.08	50.00	2.04	2.67
Maurolicus muelleri	2	2.15	2.32	4.08	0.47
Teleostei unidentified	4	4.30	0.45	8.16	0.99
Cephalopoda	10	10.75	4.53	18.37	4.74
Rossia macrosoma	5	5.38	1.18	10.20	1.72
Cephalopoda unidentified	5	5.38	3.35	10.20	2.28
Unidentified prey	20	21.51	12.57	40.82	23.48

<u>Table 2:</u> Summary of stable isotope data from white muscle tissue of *E. spinax*, from different locations, showing sampling depth (in meters), number of samples analysed (including male:female sex ratio) and individual total length (TL). Sampled depth and TL are expressed in mean (minimum-maximum values). Carbon and nitrogen ratios are given as mean (± standard deviation). All isotopic values are expressed in ‰ with δ^{13} C/ δ^{15} N representing *E. spinax* muscle isotopic composition, δ^{13} C_{phyto} and δ^{15} N_{phyto} the phytoplankton values extracted from predictive models and Δ^{13} C and Δ^{15} N the difference between shark muscle isotope values and phytoplankton modelled isotopic baselines.

Location	Depth (m)	N (M:F)	TL (cm)	δ ¹³ C	$\delta^{15}N$	$\delta^{13}C_{phyto}$	$\delta^{15} N_{\text{phyto}}$	Δ ¹³ C	$\Delta^{15}N$
Balearic Islands	589 (230-754)	33 (17:16)	26.2 (11.0-57.4)	-17.8 (0.5)	10.0 (0.4)	-27.7	4.0	9.9 (0.5)	6.0 (0.4)
Norway	243 (230-250)	51 (18:33)	43.4 (33.0-53.0)	-18.6 (0.3)	12.5 (0.7)	-25.0	0.2	6.4 (0.3)	12.3 (0.7)
Portugal	569 (490-670)	44 (19:25)	28.3 (18.6-49.0)	-18.4 (0.5)	11.1 (0.6)	-25.1 (0.6)	5.9 (0.4)	6.7 (0.7)	5.3 (0.8)
Rockall	634 (500-850)	19 (11:8)	41.6 (19.5-55.0)	-18.3 (0.5)	11.6 (0.8)	-25.2 (0.1)	0.4	7.0 (0.5)	11.2 (0.8)

Journal Pre-proof

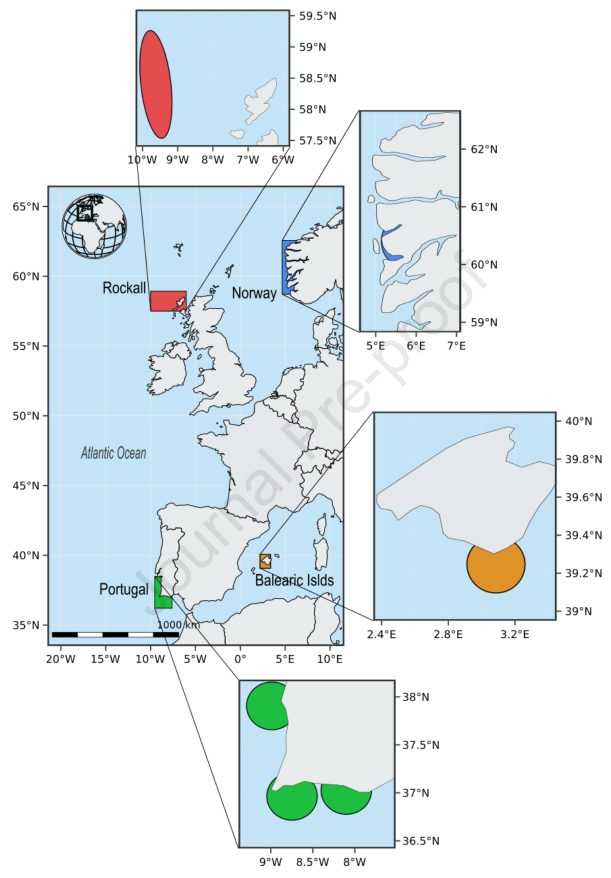
<u>Table 3:</u> Summary of Layman metrics calculated on *E. spinax* isotopic niches (Δ^{13} C/ Δ^{15} N values) and based on convex hull areas. Δ^{13} C rg and Δ^{15} N rg represent both isotopic ranges, TA the total area, CD the mean distance to centroid, NND (as mean ± standard deviation) the nearest neighbor distance (details of Layman metrics calculation can be found in Materials and Methods). All values are presented in ‰ except for TA (in ‰²).

Location	Δ ¹³ C rg	Δ ¹⁵ N rg	TA	CD	NND
Balearic Islands	2.19	1.53	1.85	0.51	0.16 ± 0.16
Norway	1.18	2.88	1.95	0.66	0.12 ± 0.07
Portugal	3.45	2.88	5.88	0.95	0.21 ± 0.14
Rockall	1.71	3.04	2.84	0.79	0.28 ± 0.21

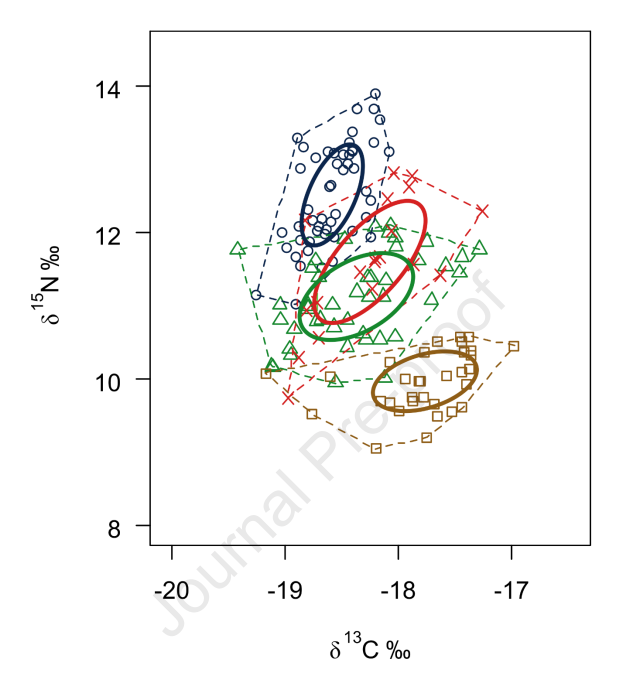
<u>Table 4:</u> Summary of *E. spinax* feeding habits. For each study, total length of sampled individuals (TL), length at which ontogenetic diet shift occurred (TL OS) and major prey before and after *E. spinax* ontogenetic diet shift (except for two studies describing constant diet) are reported. Prey groups accounting for more than 50% IRI (Index of Relative Importance) are given in bold character.

Location	TL (cm)	TL OS (cm)	Major prey Major prey before OS after OS		Publication
	10.0-49.0	~20.0	Euphausiacea (<i>M.</i> <i>norvegica</i>), Cephalopoda	Osteichthyes, Cephalopoda	Macpherson, 1980
Balearic Islands	15.0-45.0	~25.0 (*)	Natantian decapods, Euphausiacea, Cephalopoda Osteichthyes (Stomiidae, Myctophidae), Cephalopoda		Fanelli et al., 2009
	11.0-47.0	-	Cephalopoda , Te	Valls et al., 2011	
	10.2-48.3	~22.0 (**)	Cephalopoda, Telesotei, Decapods, Euphausiacea Cephalopoda, Telesotei, Decapod		Valls et al., 2017
Norway	31.0-52.0	~36.0	Euphausiacea (<i>M.</i> norvegica)	Euphausiacea (<i>M.</i> norvegica), Teleostei, Cephalopoda	This study
	11.0-33.0	11.0-33.0 - Euphausiacea (<i>M. norvegica</i>), Tele Cephalopoda			Santos & Borges, 2001
Portugal	9.1-40.1	~28.0 (***)	Euphausiacea (M. norvegica), Natantids, Teleostei	Natantids, Teleostei (Gadoids mainly), Euphausiacea, Cephalopoda	Neiva et al., 2006
Rockall	12.6-53.0	~39.0	Euphausiacea (<i>M.</i> norvegica), Teleostei (<i>M. muelleri</i>), Cephalopoda, Decapoda	Decapoda (<i>Pasiphaea</i> tarda), Cephalopoda, Other Teleostei	Mauchline & Gordon, 1983

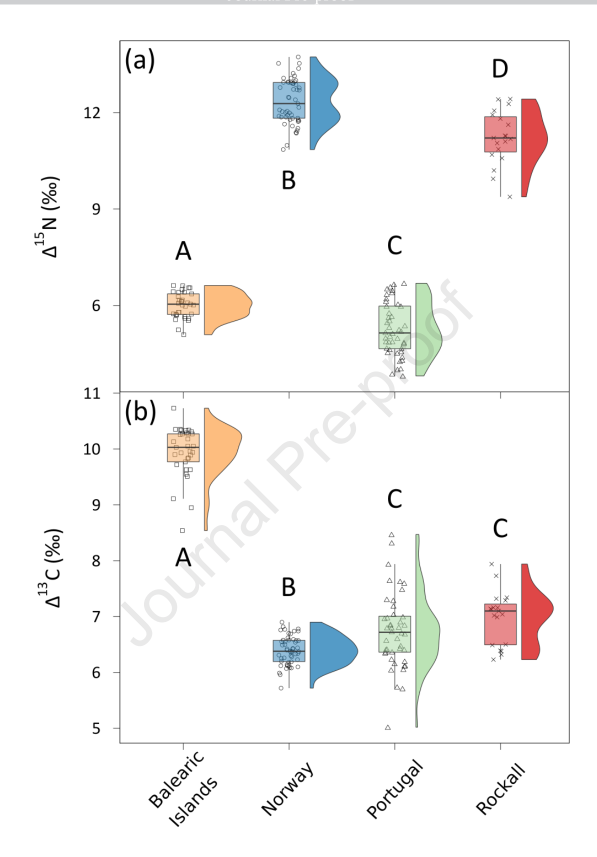
(*) Specimens < 15.0 cm presented a diet focused on Euphausiacea, fishes and to a less extend Decapoda, specimens between 15.0 and 25.0 cm had a diet mainly focused on Cephalopoda and to a less extend Decapoda and specimens > 25.0 cm focused mainly on fish. (**) OS corresponded to the absence of Euphausiacea in the diet after 22.0 cm. (***) Two OS with the first one observed around 17.0 cm when E. spinax individuals evolved from a diet focusing at 95.11 % (IRI) on Euphausiacea to a diet with 50.31 % (IRI) of Euphausiacea completed with Natantids (39.98 % IRI) and teleost fishes (9.03 % IRI). The second OS is described in the Table.



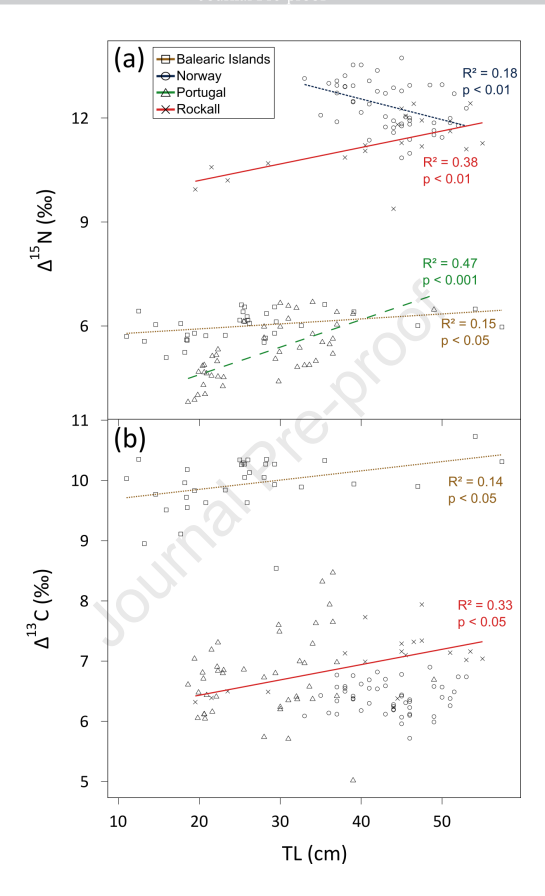
<u>Figure 1:</u> Sampling locations of velvet belly lantern sharks, *Etmopterus spinax*. Sampling area in Norway both corresponds to specimens analysed for stomach content and stable isotopes analysis.



<u>Figure 2:</u> Muscle δ^{13} C and δ^{15} N values of *E. spinax*. Brown square points correspond to the Balearic Islands, blue circle points to Norway, green triangle points to Portugal and red cross points to Rockall samples. Solid lines delimit the standard ellipse areas (SEA) and dashed lines the convex hull areas for each sampling location. The only overlapping regions between SEA occurs between Portugal and Rockall individuals (39.89%).



<u>Figure 3:</u> Boxplots of $\Delta^{15}N$ (a) and $\Delta^{13}C$ (b) values of *E. spinax* at each location. Significant differences are indicated by letters (p < 0.05).



<u>Figure 4:</u> Relationships between individual $\Delta^{15}N$ (a)/ $\Delta^{13}C$ (b) values and total length (TL) of *E. spinax*. Pearson linear regressions were applied for each location with their corresponding R² and p-value reported in the figure.

HIGHLIGHTS

- E. spinax were sampled across the northeastern Atlantic and Mediterranean Sea.
- Stomach contents suggested a similar diet.
- E. spinax fed mainly on Euphausiacea with an ontogenetic diet shift.
- Muscle carbon and nitrogen isotope compositions differed between locations.
- Isotopic differences probably resulted from mechanisms affecting the baselines.

Journal Pre-proof

Declaration of interests

☑ The authors declare that they have no known competing financial interests or personal relationships hat could have appeared to influence the work reported in this paper.
☐The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: