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# Intrinsic and extrinsic factors drive ontogeny of early-life at-sea behaviour in a marine top predator

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1 **Intrinsic and extrinsic factors drive ontogeny of**  
2 **early-life at-sea behaviour in a marine top**  
3 **predator**

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15 **ABSTRACT**

16 **Young animals must learn to forage effectively to survive the transition from parental provisioning**  
17 **to independent feeding. Rapid development of successful foraging strategies is particularly**  
18 **important for capital breeders that do not receive parental guidance after weaning. The intrinsic**  
19 **and extrinsic drivers of variation in ontogeny of foraging are poorly understood for many species.**  
20 **Grey seals (*Halichoerus grypus*) are typical capital breeders; pups are abandoned on the natal site**  
21 **after a brief suckling phase, and must develop foraging skills without external input. We collected**  
22 **location and dive data from recently-weaned grey seal pups from two regions of the United**  
23 **Kingdom (the North Sea and the Celtic and Irish Seas) using animal-borne telemetry devices during**  
24 **their first months of independence at sea. Dive duration, depth, bottom-time, and benthic diving**  
25 **increased over the first 40 days. The shape and magnitude of changes differed between regions.**  
26 **Females consistently had longer bottom-times, and in the Celtic and Irish Seas they used shallower**  
27 **water than males. Regional sex differences suggest that extrinsic factors, such as water depth,**  
28 **contribute to behavioural sexual segregation. We recommend that conservation strategies consider**  
29 **movements of young naïve animals in addition to those of adults to account for developmental**  
30 **behavioural changes.**

## 31 Introduction

32 Transition from dependence on parental provisioning to independent feeding is a critical time in the life of  
33 all animal species that receive parental care. For slow-maturing species, the first months of independent  
34 life are crucial in determining survival to recruitment, and therefore sustaining stable populations<sup>1-3</sup>.  
35 Survival depends on developing the ability to successfully find, compete for, capture and handle food  
36 resources whilst avoiding predation<sup>4,5</sup>. Juvenile behaviour, and its relationship with the development of  
37 successful feeding strategies, is receiving increasing research interest given the influence of early life  
38 survival on population dynamics<sup>1,3,6-9</sup>. Unpicking the intrinsic and extrinsic factors that affect the  
39 development of foraging skills is key to understanding population trajectories and identifying critical  
40 habitat for species during their most vulnerable life stages.

41 For air-breathing marine diving predators, such as marine mammals, sea turtles, and seabirds, the  
42 challenge of developing effective foraging strategies is particularly acute. Individuals must locate and  
43 exploit patchily-distributed prey resources in a dynamic environment, within the physiological constraints  
44 of breath-hold diving<sup>10</sup>. Studying ontogeny in wild marine predators is problematic, not least because a  
45 considerable proportion of their lives is spent at sea, often underwater, where direct observations of  
46 behaviour are difficult or impossible<sup>7</sup>. Acoustic, satellite and Global System for Mobile communication  
47 (GSM) telemetry devices have allowed ecologists to track diving predators at sea, building an increasingly  
48 clear picture of their movements and dive behaviour<sup>11,12</sup>. Logistical and practical constraints, such as high  
49 mortality rates and low re-encounter probability, mean that behavioural datasets for young animals are  
50 sparse<sup>13</sup>. Pinnipeds and seabirds are dependent on terrestrial habitat for reproduction, and young animals  
51 are large enough to carry biologging devices, therefore providing tractable opportunities to record location  
52 and behavioural data spanning the initial months of independence<sup>13</sup>.

53 Many pinniped species, including otariids and walruses (odobenids), are income breeders<sup>14</sup>: they have  
54 protracted dependency periods, during which the young learn diving and foraging skills before weaning<sup>15,16</sup>.  
55 The nursing period may last many months, or even years<sup>14</sup>. Other pinnipeds (phocids), exhibit a range of  
56 breeding strategies. Some small phocids, such as harbour seals (*Phoca vitulina*), are also income breeders,  
57 but, in contrast to otariids, pups can dive within hours of birth. Despite short dependency periods (< 1  
58 month<sup>17</sup>), harbour seal pups can develop diving skills during suckling and may accompany their mothers  
59 on foraging excursions<sup>18</sup>. Synchronous diving of mothers and pups during lactation also occurs in some ice-  
60 breeding phocids<sup>19</sup>. Larger phocid species, such as elephant (*Mirounga spp.*), hooded (*Cystophora cristata*)  
61 and grey (*Halichoerus grypus*) seals, are usually capital breeders, and pups are abruptly abandoned at the  
62 natal site after a brief nursing period<sup>14</sup>. Grey seals, for example, suckle for 15-21 days<sup>20</sup>. Pups then undergo  
63 a post-weaning fast, usually on land, of between nine and 40 days, during which time they lose up to 25%  
64 of their body mass<sup>21,22</sup>. After departure from the natal colony, they must learn to dive and find food without  
65 maternal provisioning, or the benefit of observing the foraging behaviour of their mother<sup>23</sup>. Furthermore,  
66 they must do this before their remaining blubber and protein reserves are depleted to critical levels and  
67 terminal starvation begins<sup>24</sup>.

68 Swimming in cold water and diving to depth is energetically costly, and seal pups have a higher surface  
69 area to volume ratio, higher mass-specific metabolic rate and lower mass-specific oxygen storage capacity  
70 than adults<sup>25,26</sup>. In contrast to adults, young seals repeatedly dive up to their physiological limits and  
71 foraging efficiency is therefore lower because they must spend longer at the surface to recover<sup>15,27</sup>.  
72 Maximum diving capability increases in grey seal pups during the first months at sea<sup>24</sup>, but little is known  
73 about the development of their routine behaviours. First year mortality is high and variable between years  
74 for grey seals<sup>28-30</sup>, which has a profound effect on population dynamics<sup>31</sup>. Moreover, first year survival  
75 probability appears to be three times greater for females than males, regardless of body condition at  
76 weaning<sup>29</sup>. Differences in survival between male and female pups could be linked to development of sex-  
77 specific diving behaviour, leading to the sex difference in foraging strategies underpinned by sexual size  
78 dimorphism in adults<sup>32</sup>. In general, adult grey seals make repeated, short duration (3-10 days) foraging  
79 trips offshore within shelf seas, diving to the bottom to exploit benthic and demersal prey, and returning  
80 to coastal 'haul-out' sites<sup>33,34</sup>. Most research has focussed on adult movements and foraging strategies.  
81 Whilst some work has investigated foraging in grey seal juveniles and YOY<sup>35,36</sup>, and others have studied pup  
82 behaviour on and around the colony<sup>37,38</sup>, only Bennett *et al.*<sup>24</sup> have examined the ontogeny of at-sea  
83 behaviour in recently-weaned pups across their first months of nutritional independence. Previous studies  
84 have demonstrated sex differences in the foraging behaviour of grey seal adults (seals of reproductive  
85 age)<sup>32,36</sup>, juveniles (> 12 months old)<sup>36</sup>, and young-of-the-year (YOY; 5 months old)<sup>35</sup>. Sex differences in  
86 behaviour thus emerge from an early age<sup>35</sup>, but the timing of their onset is unknown. Development of diving  
87 and learning of successful foraging behaviour is also likely to be shaped by local experience, and the  
88 environment that pups encounter when they first go to sea. Oceanographic conditions and prey availability  
89 vary among regions, presenting different challenges for different subpopulations. Together, these factors  
90 may confer regional differences in the ontogeny of diving behaviour and thus the development of successful  
91 foraging strategies for grey seal pups.

92 The United Kingdom (UK) is home to ~38% of the world grey seal population<sup>39</sup> and has an obligation under  
93 European Union (EU) legislation to maintain this population in favourable conservation status (FCS)<sup>40</sup>. As  
94 part of this obligation, critical habitat must be identified for this species both on land and at sea to assess  
95 and mitigate anthropogenic disturbance. Current UK conservation management for grey seals at sea is  
96 largely based upon observations of adult movement<sup>39</sup>. Foraging behaviour has not yet been described for  
97 grey seal pups, however, given that they undergo profound physiological development during their initial  
98 months of independent life<sup>24,25</sup>, coupled with a need to explore their environment and develop knowledge  
99 of potential foraging areas, we should not expect their behaviour and habitat requirements to be the same  
100 as for adults. As pups develop diving skills, grow larger and acquire knowledge of their surroundings, we  
101 might expect that their behaviour begins to converge on that of adults, since adult behaviour represents  
102 successful foraging patterns. The main aim of this study, therefore, was not to quantify foraging in grey seal  
103 pups, but to investigate changes in at-sea behaviours relevant to the development of successful foraging  
104 skills during their first four months of independent life at sea. We used a unique, large ( $n = 52$  individuals)  
105 animal-borne satellite and GSM telemetry dataset of location and dive (time-depth) data from recently-  
106 weaned pups born at six different colonies around the UK (Table 1). Ontogeny of foraging behaviour has

107 been characterised in young seals by reductions in trip metrics (duration and distance), and increases in  
108 dive metrics (depth, duration, proportion of dives that are benthic, bottom time and proportion of day spent  
109 diving) with age<sup>15,24,41,42</sup>. Such changes in these metrics are indicative of an individual's ability to maximise  
110 foraging opportunities within individual dives and/or over foraging trips, and are thus representative of  
111 greater foraging efficiency<sup>15,24,41,42</sup>. Thus, using generalized estimating equations in a generalized additive  
112 model framework (GEE-GAM), we investigated how these variables changed over time and compared the  
113 trajectories between the sexes and two distinct geographic regions (Celtic and Irish Seas (hereafter CIS)  
114 and North Sea (hereafter NS); Fig. 1). Furthermore, sexual segregation of foraging habitat may be  
115 manifested in the depth of water where males and females dive<sup>35</sup>. We therefore examined differences in  
116 the bathymetric depth of dive locations in the same way.

117

## 118 **Results**

### 119 **Trip behaviour**

120 All pups remained within the limits of the continental shelf, but NS individuals had a much wider dispersal  
121 pattern, and several pups travelled along the shelf break (Fig. 1). Although NS pups travelled far from their  
122 natal colonies on individual trips, all returned to haul-out locations on the east coast of Scotland and  
123 England. No pup crossed the shelf break into waters > 200 m deep. However, one male from the Isle of May  
124 travelled between the UK and Norway on multiple occasions, diving to the bottom of the Norwegian Trench  
125 (Fig. 1b; > 200 m). In general, pups from both regions explored new areas before settling into repeated trip  
126 behaviour, hauling-out in one or more locations and commuting back and forth to foraging grounds, as  
127 observed in adults<sup>34</sup> (Fig. 2). Many NS pups undertook a prolonged exploratory phase shortly after leaving  
128 the colony, with 69% of pups (n=24) spending > 20 days offshore without returning to the coast, and some  
129 individuals exceeding 60 days offshore, which is substantially greater than typical trip durations seen in  
130 adults<sup>34</sup>. Only 18% of CIS pups (n=3) performed a trip with duration > 20 days. CIS pups remained much  
131 closer to land, generally dispersing along the coast of Wales and the Republic of Ireland (Fig. 1a). One female  
132 travelled south towards the north coast of France before returning to the south coast of England. Some  
133 individuals made repeated trips into the middle of the Celtic Sea, while others remained within 30 km of  
134 the coastline, and never travelled > 50 km from their natal colony (Fig. 3).

135 Trip duration increased significantly with time after leaving the colony for pups from both regions (Table  
136 2; GEE-GAM;  $\chi^2_3 = 15.2$ ,  $p = 0.002$ ), peaking at around 70 days before declining (Fig. 4a-b). However, trip  
137 duration was significantly longer for NS pups than CIS pups (Fig. 4a-b; GEE-GAM;  $\chi^2_1 = 66.1$ ,  $p < 0.001$ ).  
138 There was no significant difference in trip duration between males and females in either region (GEE-GAM;  
139  $\chi^2_1 = 1.4$ ,  $p = 0.233$ ). Trip distance was also significantly affected by time since departure for pups from both  
140 regions (Table 2; GEE-GAM;  $\chi^2_3 = 8.2$ ,  $p = 0.042$ ), peaking at around 70 days then declining (Fig. 4c-d).  
141 However, there was a significant effect of an interaction between region and sex on trip duration (Fig 4d;  
142 GEE-GAM;  $\chi^2_1 = 4.73$ ,  $p = 0.03$ ); NS pups travelled consistently further than CIS pups. CIS males travelled  
143 further than females, whilst there was no obvious sex difference in trip distance for NS pups.

144

## 145 **Dive behaviour**

146 A three-way interaction between time since departure, region and sex best explained variation in daily  
147 mean maximum dive depth (Table 2; GEE-GAM;  $\chi^2_3 = 13.6$ ,  $p = 0.003$ ). Pups increased their dive depth  
148 rapidly over the first 40 days, except for CIS females, which showed a prolonged, more moderate increase  
149 (Fig. 5a-b). Sex differences in the change in dive depth over time were apparent in CIS pups, with males  
150 diving significantly deeper than females from 20-60 days after leaving the colony (Fig. 5b). The population  
151 mean maximum depth for CIS males during this period reached  $\sim 40$  m, whilst females achieved  $\sim 25$  m.  
152 Throughout the time series, NS pups dived significantly deeper than CIS pups, with both males and females  
153 reaching a maximum daily mean of  $\sim 50$  m.

154 A three-way interaction between time since departure, region and sex best explained variation in daily  
155 mean bathymetric depth of dive locations (Table 2; GEE-GAM;  $\chi^2_3 = 10.4$ ,  $p = 0.016$ ). NS pups and CIS males  
156 dived in increasingly deep water over the first 40 days after departure from the colony (Fig. 5c). CIS females  
157 remained in shallower water than males throughout the first four months at sea, averaging depths of  $\sim 30$   
158 m whilst mean bathymetric depth for male dives reached up to  $\sim 60$  m (Fig. 5d). No significant sex difference  
159 was evident in bathymetric depth of dive locations for NS pups. Both male and female NS pups dived in  
160 significantly deeper water than CIS pups, reaching a maximum daily mean of  $\sim 80$  m.

161 The daily mean proportion of dives that were benthic changed with time since departure, and the dynamic  
162 of this change was different between the regions (Table 2; GEE-GAM;  $\chi^2_3 = 13.1$ ,  $p = 0.004$ ). Pups from both  
163 regions increased the proportion of benthic dives rapidly over the initial 40 days. This reached an  
164 asymptote for NS pups (Fig. 5e), but continued to increase for CIS pups (Fig. 5f). The trend showed some  
165 evidence of a decline in the latter half of the time series for NS pups, but confidence intervals were wide  
166 (Fig. 5e). Females performed a greater proportion of benthic dives than males throughout the time series  
167 in both regions (GEE-GAM;  $\chi^2_1 = 5.2$ ,  $p = 0.023$ ). The daily mean proportion of benthic dives reached a peak  
168 at  $\sim 0.5$  for NS females,  $\sim 0.6$  for CIS females,  $\sim 0.4$  for NS males, and  $\sim 0.5$  for CIS males. Confidence intervals  
169 for the sexes overlapped in both regions. The effect of bathymetric depth on the proportion of dives that  
170 were benthic is presented in Supplementary Information (Supplementary Results: Effects of bathymetric  
171 depth on benthic diving).

172 Daily mean dive duration was best explained by an interaction between time since departure and region  
173 (Table 2; GEE-GAM;  $\chi^2_3 = 16.4$ ,  $p < 0.001$ ). There was no significant effect of sex on this metric (GEE-GAM;  
174  $\chi^2_1 = 2.5$ ,  $p = 0.117$ ). Similar to dive depth and bathymetric depth, pup dive duration increased rapidly over  
175 the initial 40 days at sea for both regions, before declining over the following 60 days (Fig. 6a-b). Peak mean  
176 dive duration for NS pups was marginally longer than for CIS pups (NS:  $\sim 140$  s, CIS  $\sim 130$  s).

177 Temporal patterns in bottom time differed between regions (Table 2; GEE-GAM;  $\chi^2_3 = 14.9$ ,  $p = 0.002$ ). CIS  
178 pups showed a strong increase in bottom time over the initial 40 days at sea, before levelling off, then a  
179 further increase at  $\sim 100$  days. NS pups showed a moderate increase over the whole time series, with

180 bottom time remaining between 40-50% of dive duration (Fig. 6c-d). In addition, sex differences in bottom  
181 time differed between the regions (GEE-GAM;  $\chi^2_1 = 9.3$ ,  $p = 0.002$ ). In both regions, females achieved higher  
182 bottom times than males (although 95% confidence intervals overlapped for NS pups; Fig. 6c). The  
183 difference between males and females was more pronounced in CIS pups: females achieved a maximum  
184 mean of ~70% of the dive spent in the bottom phase, whilst males achieved a maximum mean of ~55%  
185 (Fig. 6c-d).

186 Time spent diving per day varied significantly with time since departure, and the shape of this relationship  
187 was affected by sex (Table 2; GEE-GAM;  $\chi^2_3 = 13.9$ ,  $p = 0.003$ ) and region (GEE-GAM;  $\chi^2_3 = 15$ ,  $p = 0.002$ ).  
188 The sex difference was comparable between both regions (GEE-GAM;  $\chi^2_1 = 0.02$ ,  $p = 0.885$ ). NS pups began  
189 diving ~14 hrs per day, then reduced time spent diving in the third month to ~10 hrs for males and ~12  
190 hrs for females (Fig. 6e). CIS females initially spent ~11 hrs diving per day, which rose steadily to ~13 hrs  
191 in the third month (Fig. 6f). CIS males initially spent ~10 hrs per day diving, which rose steeply to ~13 hrs  
192 in the first month before declining back to ~11 hrs in the third month (Fig. 6f).

193

## 194 Discussion

195 This study reveals that sexual segregation of behaviour can be exhibited as early as nutritional  
196 independence in capital breeders. Female pups from both regions spent more time diving per day than  
197 males. CIS females made shorter distance trips than males, diving in shallower water and achieving a higher  
198 proportion of the dive duration in the bottom phase. The same level of sexual segregation in depth,  
199 proportion bottom time and trip duration was not observed in NS pups, suggesting that sex differences in  
200 the ontogeny of foraging behaviour may be mediated by extrinsic factors. In both regions, pup behaviour  
201 changed rapidly: dive duration, depth, bottom time and benthic diving increased over the first 40 days after  
202 leaving the colony. These findings are important in the context of both foraging ecology and conservation  
203 management, as we outline below.

204 Grey seal adults exhibit substantial sexual size dimorphism<sup>23</sup>, which is thought to drive differences in  
205 feeding areas<sup>32</sup>. Grey seal pups are not size-dimorphic<sup>24</sup>, but seal pups and juveniles may experience  
206 differences in energy requirements before overt size and body composition differences emerge<sup>43</sup>. Kelso *et*  
207 *al.*<sup>43</sup> found that male northern elephant seal (*M. angustirostris*) pups had higher rates of energy expenditure  
208 than females during the post-weaning fast, but were more effective at sparing protein reserves. These  
209 differences are likely related to the development of sex-specific metabolic strategies required for successful  
210 breeding<sup>43</sup>. Differences in metabolic demand during the ontogeny of foraging behaviour could therefore  
211 drive sex-specific feeding strategies and habitat requirements. Our findings support this possibility; we  
212 found that females from both regions spent longer performing behaviours consistent with foraging across  
213 two different temporal scales (individual dives and 24 h period). At the individual dive scale, time in the  
214 bottom phase is indicative of time at potential foraging depth, with the descent and ascent phases of the  
215 dive representing the transit to and from any potential prey patch<sup>10</sup>. Despite the lack of sex difference in  
216 total dive duration, females spent longer in the bottom phase than males relative to total dive duration. At

217 the 24 h scale, females spent on average 2 h more diving than males in both regions. We also found a  
218 moderate sex difference in the proportion of dives that reached the seabed, with females performing more  
219 benthic dives than males. Females may therefore have increased chance of prey capture during individual  
220 dives, which could represent an energetic advantage<sup>10,44</sup>, and contribute to higher survival probability of  
221 female pups<sup>29</sup>. However, we cannot exclude the possibility that females spend more time diving because  
222 they are searching and are unsuccessful. Using direct observations of prey capture (i.e. stomach  
223 temperature telemetry, accelerometers or video cameras) to ground-truth putative foraging as identified  
224 from location and dive data would help to evaluate foraging success<sup>45</sup>, and draw links between differences  
225 in ontogeny of foraging behaviour and survival probability.

226 Sex differences in bottom time, proportion of benthic dives, and time spent diving per day may be related  
227 to differences in the type and quality of prey items consumed by male and female pups. For example, if  
228 females target lower energy prey items, they will need to spend longer foraging than males for the same  
229 energetic gain. Grey seals adults are benthic foragers with a broad diet that varies between the sexes<sup>46,47</sup>.  
230 Beck *et al.*<sup>47</sup> used quantitative fatty acid analysis to investigate niche breadth in grey seals in the northwest  
231 Atlantic. They found that the diet composition of YOY animals was significantly broader than that of adults,  
232 but found no sex differences for young animals. However, grey seal diet varies regionally and seasonally<sup>46,47</sup>,  
233 and therefore extrinsic factors unique to certain locations may shape sex differences in diet for young  
234 animals. No specific information currently exists on the diet of recently-weaned grey seal pups in the UK  
235 once they have left the colony due to the logistical constraints of collecting tissue and/or faecal samples  
236 specifically from this age-class. However, a recent study of stable isotope ratios obtained from the teeth of  
237 older juvenile grey seals in the North Sea suggests that they feed on a wide variety of low trophic level,  
238 benthic prey close to shore<sup>48</sup>. The sharp increase in proportion of benthic dives over the first 40 days, and  
239 the subsequent reduction in trip distance, may therefore be indicative of pups learning to exploit benthic  
240 prey, and finding foraging grounds closer to shore where they can effectively reach the bottom. Additional  
241 dive analysis also suggests that shallow waters < 20 m deep may represent important foraging habitat for  
242 grey seal pups (see Supplementary Results: Effects of bathymetric depth on benthic diving).

243 Water depth is an important regulating factor in foraging behaviour and habitat preference in older grey  
244 seals<sup>49,50</sup>. Breed *et al.*<sup>35</sup> reported that adult and YOY females in the northwest Atlantic population forage in  
245 shallower water than males. Our data from CIS pups, showing that females dived in significantly shallower  
246 water than males, support these findings and suggest that water depth may play a key role in the  
247 development of habitat (and possibly diet) segregation among the sexes in some regions. We also found a  
248 moderate sex difference in trip distance for CIS pups, with males travelling further than females. Given that  
249 there was no sex difference in trip duration, this may mean that CIS males travel further offshore to forage  
250 compared to females, accessing deeper water, and potentially spending longer travelling per unit time  
251 spent foraging than females. CIS females performed a greater proportion of benthic dives in shallow water  
252 (<20 m) than males (see Supplementary Results: Effects of bathymetric depth on benthic diving). The fact  
253 that CIS pups dived in shallower water than NS pups likely means that they were able to achieve greater  
254 dive bottom time and proportion of benthic dives as they spent less time in the ascent and descent phases

255 of the dive. Sex differences in trip distance and water depth of dive locations were not strongly evident for  
256 NS pups. As with other metrics, sex differences may be mediated by extrinsic factors that vary among  
257 regions, such as prey distribution, physical oceanography, and the diversity of available habitats. In general,  
258 the North Sea is a more homogeneous ecosystem, with less variation in bathymetry and habitat types than  
259 the Celtic and Irish Seas<sup>51</sup>, which may reduce sexual niche separation in NS pups.

260 Intra and inter-specific competition may impact trip distance and duration in central place foragers.  
261 Juvenile grey seals in the northwest Atlantic travel further and for longer on foraging trips than adults,  
262 likely as a result of competitive exclusion from the best foraging grounds closer to shore<sup>52</sup>. Age-related  
263 segregation has also been reported for other phocid species<sup>53</sup>. We found that NS pups travelled further  
264 offshore and performed longer trips than CIS individuals. Population density of grey seal adults is much  
265 higher on the east coast of Scotland compared to the Celtic and Irish Seas<sup>39,54</sup>. Moreover, Russell *et al.*<sup>36</sup>  
266 showed that adult males in the North Sea reduce their time spent travelling to foraging locations in winter,  
267 whilst juveniles show an increase. Given that NS pups leave the colony during the winter months, and we  
268 see the longest trips performed during this time, competitive exclusion by conspecifics may be a feature of  
269 movement patterns specifically during the winter, forcing pups to make longer trips further offshore. In  
270 addition, harbour seals are present in coastal regions of the North Sea, but not in the Celtic and Irish Seas<sup>54</sup>.  
271 Inter-specific competition may also contribute to NS pups travelling further offshore than CIS pups.

272 Our results show that NS pups can make trips of over two months in duration, travelling greater distances  
273 than commonly observed in adult foraging trips and hauling out less frequently<sup>34</sup>. We also found that pups  
274 significantly reduced their trip duration and distance in the third month (Fig. 2). A similar temporal  
275 dynamic has been observed in other phocids, with young seals reducing trip duration after an initial  
276 increase<sup>41</sup>, and may be indicative of an increase in foraging efficiency, or a change in foraging strategy as  
277 pups age. Moreover, the higher initial trip duration and distance may represent an exploration phase in the  
278 development of NS pups. Votier *et al.*<sup>55</sup> found that immature northern gannets (*Morus bassanus*) develop  
279 knowledge of foraging grounds during early-life exploratory trips. This may also be the case for grey seal  
280 pups, as, like gannets, they receive no parental guidance in the location of foraging resources. Furthermore,  
281 we found that some pups returned to forage repeatedly in areas that they had previously discovered during  
282 their initial exploratory trip (Fig. 2). Exploration may therefore be an important behaviour in determining  
283 future foraging success<sup>8</sup>.

284 CIS pups also performed exploratory trips, although their duration and distance was lower than those  
285 performed by NS pups. Individuals from NS colonies are not as geographically constrained as CIS pups by  
286 the proximity of land and shelf edge and therefore have more marine space to explore. Upon leaving the  
287 colony, CIS pups are more likely to encounter coastline, and therefore suitable haul-out locations, than pups  
288 in the North Sea. Alternatively, the offshore phase could be driven by environmental variables not  
289 measured in this study. For example, tidal currents may direct pups further from land in the North Sea. The  
290 reduction in trip distance after 60 days for NS pups may therefore be related to a seasonal change in  
291 physical oceanography, or an increase in their ability to resist surface currents as muscle strength improves.

292 Grey seals are known to rest at sea<sup>36</sup>, and this study provides further evidence that they do not need to  
293 return to shore to rest, even when very young.

294 Our results show that pup movements can change rapidly throughout the initial months at sea. Therefore,  
295 accurately quantifying foraging effort from these data may require extension of current analytical  
296 techniques, such as state-space models (SSMs)<sup>45</sup>, to account for temporal changes in movement patterns.  
297 Moreover, as a priority for future work, analysis of pup foraging habitat preference may allow us to infer  
298 potential prey species based on habitat features such as substrate type, and further assess the implications  
299 of early-life sexual segregation in movement patterns for foraging ecology.

300 In addition to ontogenetic changes in muscular and cardio-vascular systems, oxygen storage capacity and  
301 metabolic rate, and the development of knowledge of profitable foraging areas, there are likely to be  
302 seasonal changes in foraging habitat and prey distribution which may further explain differences in pup  
303 behaviour over time. Given that pups leave the colony on different dates in both regions (see  
304 Supplementary Note: Colony departure dates), local conditions may dictate some of the patterns observed  
305 here. Bennett *et al.*<sup>56</sup> have shown that maximum dive depth of adult southern elephant seals may be  
306 regulated by seasonally-mediated factors, however, due to a paucity of tracking data from post-breeding  
307 adult grey seals in the UK, seasonal changes in at-sea behaviour are unclear. It was therefore not possible  
308 to disentangle ontogeny from seasonal effects on pup behaviour. Furthermore, some of the variance in  
309 early-life behavioural ontogeny may be explained by the fact that post-weaning fast duration varies among  
310 individuals<sup>24</sup>, and age at the point of departure from the colony is not equal for all pups. Natal and weaning  
311 dates were not known for all pups in this study, and time since departing colony was therefore used as a  
312 measure of at-sea experience. Future research should aim to achieve simultaneous tagging of adults,  
313 juveniles and pups, coupled with colony-based monitoring, which will allow us to further tease apart  
314 intrinsic and extrinsic drivers of variation in grey seal foraging behaviour and investigate the potential for  
315 competitive exclusion<sup>36</sup>.

316 Investigating the factors that affect the ontogeny of early-life behaviours is key to understanding how  
317 populations may respond to natural and anthropogenic threats. Bennett *et al.*<sup>22</sup> suggested that grey seal  
318 pups have an average of 36 days in which to find food after leaving the colony before their protein reserves  
319 are critically depleted and starvation occurs. Our results show that profound changes in pup behaviour  
320 happen during the first 40 days after departure from the colony, indicating this initial period at sea is likely  
321 of particular importance for development of effective foraging strategies. Consequently, pups may be most  
322 vulnerable to disturbance from a number of growing anthropogenic activities, such as increased vessel  
323 traffic<sup>57</sup>, intensive fishing practices<sup>58</sup> and offshore construction<sup>59</sup> during this period, with substantial  
324 consequences for survival. Given the importance of early-life survival for maintaining stable populations<sup>31</sup>,  
325 and the rapid development of key behaviours during this period, conservation managers should make  
326 special considerations for pups during their initial months at sea to effectively mitigate these threats and  
327 avoid population-level impacts. With continuing development of biologging technology and analytical  
328 techniques, further work is urgently needed to fully explore and describe the ontogeny of fundamental

329 behaviours in naïve marine predators and identify critical habitat for young animals during their most  
330 vulnerable life stage.

331

## 332 **Methods**

### 333 **Instrumentation**

334 Two different telemetry device models were deployed on 52 recently-weaned grey seal pups at six UK  
335 breeding sites in 2001 and 2002<sup>24</sup>, and in 2009 and 2010<sup>60</sup> (Table 1). Earlier deployments (2001-2002;  
336 n=21) were Argos Satellite Relay Data Loggers (SRDL; Sea Mammal Research Unit, UK), and later  
337 deployments (n=31) were Fastloc® GPS-GSM tags (GPS phone tags; Sea Mammal Research Unit, UK).  
338 Individuals were captured post-weaning for device application. When anaesthesia was required (due to  
339 additional procedures not related to this study; CIS 2010 and all Isle of May deployments), pups were  
340 administered with 0.025 mg kg<sup>-1</sup> intravenous Zoletil<sub>100</sub>® (Virbac, France)<sup>24,60</sup>. Following McConnell *et al.*<sup>34</sup>,  
341 a tag was glued to cleaned, dried fur at the base of the skull using RS Quick-Set Epoxy Adhesive (RS  
342 Components Ltd., UK; 2001-2009), or Loctite® 422™ cyanoacrylate adhesive (Henkel, UK; 2010). All  
343 experimental protocols were carried out with UK Home Office approval under project licences #60/2589  
344 (2001-2002), and #60/4009 (2009-2010), in accordance with the Animals (Scientific Procedures) Act 1986.  
345 In total, 7057 days of data were recorded from 52 pups (for information on tag duration see Supplementary  
346 Note: Tag duration).

347

### 348 **Telemetry data processing**

#### 349 ***Horizontal movement data***

350 Whilst both SRDL and GPS-GSM devices transmitted location data at irregular intervals, mean number of  
351 location fixes achieved per day was much higher for GPS-GSM tags (Table 1). Argos-derived location  
352 estimates from SRDLs also carry a greater spatial error, ranging from 50 m to > 2.5 km<sup>61</sup>. Erroneous Argos  
353 location observations were eliminated using the standard technique of filtering with a maximum speed  
354 threshold of 2 ms<sup>-1</sup>.<sup>62</sup> Remaining locations were then processed with a Kalman filter to improve location  
355 accuracy<sup>54</sup>. Kalman filter observation model parameters were taken from Vincent *et al.*<sup>61</sup>, and process  
356 model parameters were based on average speeds of 142 seal GPS tracks<sup>54</sup>. Erroneous GPS locations were  
357 identified and excluded using residual error thresholds and number of satellites<sup>36</sup>.

358 Devices also recorded dive and haul-out data derived from integrated conductivity and pressure sensors.  
359 Following Russell *et al.*<sup>36</sup>, a seal's location during a haul-out event was taken as the mean of all latitude and  
360 longitude estimates during the time hauled-out. If no location estimates were recorded during the haul-out  
361 interval, the location was derived using linear interpolation to a midpoint between the pre and post  
362 observed location fixes. Interpolated haul-out locations were flagged as unreliable if there was no adjacent

363 observed location within 6 h. The location data were then restricted to discrete ‘trips’ between haul-out  
364 events. Trips were only included in the analysis if they had a reliable haul-out location on land at both the  
365 beginning and end. One individual hauled-out repeatedly on an offshore oil rig in the central North Sea >  
366 250 km from land; these haul-outs were classed as on land and the associated trips were included in the  
367 analysis. Seals often wait in the water between haul-out events for tidal sites to become available, when  
368 they may sleep either on the seabed or at the surface<sup>63</sup>. To exclude this behaviour, as it is not relevant to  
369 foraging, trips < 8 hr in duration and with a maximum distance < 500 m from the coast were also omitted  
370 from the analysis<sup>64</sup>. Finally, as tag duration varied between individuals (from 13 to 337 days; see  
371 Supplementary Note: Tag duration), data were clipped at 120 days after leaving the colony to ensure a  
372 robust sample size throughout the time series for statistical analysis<sup>65</sup>. Sample sizes are presented  
373 alongside rug plots (Fig. 4). The resulting dataset comprised location and haul-out data from 52 individuals;  
374 23 males and 29 females (Table 1; Fig. 1; 836 trips). The duration and total distance of each trip was  
375 calculated alongside days since first leaving the natal colony at the mid-point of the trip. Total distance was  
376 calculated as the sum of all step lengths between successive location fixes during a trip, regularised to 30  
377 min intervals. Days since leaving colony was used to give a measure of the at-sea experience of the pup.

378

#### 379 *Dive data*

380 GPS-GSM tags classified dives as periods when the pressure sensor recorded depths > 1.5 m for > 8 s. These  
381 devices recorded depth readings at 4 s intervals throughout a dive, which were then abstracted to 11  
382 inflection points by an algorithm onboard the device before data transmission<sup>66</sup>. Although SRDLs also  
383 recorded dive data, tag parametrisation was different to that of GPS-GSM tags (SRDLs only recorded dives >  
384 6 m depth with four inflection points). Furthermore, the lower frequency of successful transmissions and  
385 higher spatial error of concomitant Argos-derived location estimates meant that SRDL dive data could not  
386 be accurately matched to a location, and thus to bathymetric depth, and were therefore excluded from all  
387 dive analyses. For GPS-GSM dive data, the maximum dive depth and total dive duration were extracted for  
388 each dive. A dive was treated as any time below the depth threshold (1.5 m). Devices also transmitted two-  
389 hourly summaries of data, detailing the proportion of time the device was in either “haul-out”, “dive” or  
390 “cruise” (device is wet and above 1.5 m) mode. These data were used to calculate the total number of hours  
391 spent diving per individual per day. Only days with data for all twelve summary intervals were used.

392 To investigate changes in the proportion of benthic dives performed by pups, and the bathymetric depth of  
393 dive locations, dives were first matched to adjacent location fixes in time using the mid-point between dive  
394 start and end times. The location for each dive mid-point was then calculated using linear interpolation  
395 between prior and post location fixes. Interpolated dive locations with no adjacent observed location fix  
396 within 15 min could not be accurately matched with bathymetric depth data and were therefore excluded  
397 from the analysis. Bathymetric depth was extracted for each dive location from the harmonised 1/8 arc  
398 minute \* 1/8 arc minute (~230 m) gridded Digital Terrain Model (DTM) for European Waters which is  
399 freely-available through the European Marine Observation and Data Network (EMODnet) Portal for

400 Bathymetry<sup>67</sup>. Benthic dives were classified following Ramasco *et al.*<sup>68</sup>, using a mixture distribution model  
401 approach (see Supplementary Methods: Classification of benthic dives). The bathymetric depth range of  
402 the study area is shown in Fig. 1. After filtering, a number of dives (15%) recorded a null or positive  
403 bathymetric depth value, due to interpolated dive locations falling too close to the coast, and were  
404 subsequently removed. As with trip data, the resulting dive dataset was clipped to 120 days after leaving  
405 the colony to ensure a robust sample size throughout the time series. Sample sizes are presented alongside  
406 rug plots (Figs. 5-6). Lastly, as seals may perform successive shallow dives while resting close to haul-outs,  
407 and this is not related to foraging behaviour, any dive with a maximum depth < 5 m was excluded. The final  
408 dataset comprised 102,800 dives from 31 individuals (Table 1).

409

### 410 **Statistical analysis**

411 Trip (duration and distance) and dive metrics (depth, bathymetric depth of dive locations, proportion of  
412 dives that were benthic, duration, bottom time and proportion of day spent diving) were analysed using  
413 generalised estimating equations within a generalized additive model framework (GEE-GAMs) using the  
414 “geepack” and “splines” packages<sup>69</sup> in R<sup>70</sup>. The GAM approach allows the inclusion of smoothed terms to  
415 investigate non-linear relationships<sup>71</sup>. However, GAMs are not robust to the serial autocorrelation within  
416 individuals that is inherent in longitudinal telemetry datasets. GAMs can be extended to include  
417 autocorrelation structures and random effects, however the GEE approach allows the inclusion of an  
418 unstructured correlation coefficient, which is more appropriate for telemetry data as it estimates all  
419 correlations between within-individual observations independently<sup>72</sup>. Furthermore, this method allows  
420 the prediction of population mean responses by averaging across individuals. This approach has been  
421 previously applied to study temporal movement trends in seal telemetry datasets<sup>36</sup>.

422 We investigated ontogeny in pup dive behaviour using a number of metrics, at a temporal resolution of one  
423 day by calculating daily means per individual per day. As pups grow, their muscular and cardio-vascular  
424 systems develop, and they convert blubber into lean mass, becoming less buoyant<sup>25,73</sup>. Their ability to dive  
425 to, and remain at depth should therefore increase over time<sup>24</sup>. Daily mean dive maximum depth and  
426 duration were used to track changes in diving ability over time. For air-breathing benthic foragers, the  
427 depth of water in which dives occur is also relevant to their ability to dive to and remain at foraging depth.  
428 The bathymetric depth of water where dives occurred was also modelled in the same way. Optimal diving  
429 theory (ODT) suggests that benthic foragers will maximise time at the seabed (and therefore probability of  
430 successful foraging), and minimise time spent in the ascent and descent phases of a dive and at the surface<sup>74</sup>.  
431 We therefore investigated changes in the proportion of dives that were benthic, and in dive bottom time  
432 (the proportion of a dive’s duration spent at > 80% of the maximum dive depth; a measure of time spent at  
433 foraging depth relative to descent and ascent phases of a dive)<sup>75</sup>. Lastly, pups may maximise time spent  
434 underwater (and therefore foraging opportunities) over bouts of short dives, rather than individual long  
435 dives<sup>10,44</sup>. We therefore investigated changes in the mean proportion of time spent diving per individual  
436 per day (24 h period).

437 Pup behaviour may change through time, and the dynamics of this change may differ between the sexes,  
438 and/or geographic regions (due to differences in habitat features such as coastal geography, prey  
439 availability and bathymetry). Therefore, response variables were analysed in separate models as a function  
440 of time since leaving colony (days; as a smoothed term), sex (as a categorical term) and region (as a  
441 categorical term) in a three-way interaction. Model selection was performed by backwards hypothesis  
442 testing from GEE-based p-values until arriving at a minimum adequate model. Colonies were assigned to  
443 one of two geographic regions (Table 1; North Sea or Celtic and Irish Seas). There was considerable spatial  
444 overlap of areas used by pups from colonies within each of the two wider geographic regions (Fig. 1), such  
445 that region rather than colony was used in the models for the sake of parsimony, and to maximise statistical  
446 power. 95% confidence intervals around model-predicted means were calculated by parametric  
447 bootstrapping using GEE-based uncertainty parameters<sup>54</sup>. Scale-corrected Pearson's residuals were  
448 checked for normal distribution by visual inspection in all models. For models with continuous response  
449 variables (all except bottom time, benthic diving and proportion of day spent diving), Gamma and Poisson  
450 error structures were considered to improve normality, but in all cases a Gaussian error structure with log-  
451 link function proved superior. Bottom time, proportion of dives that were benthic and proportion of day  
452 spent diving (proportion data) were modelled with a binomial error structure with logit-link function.

453

#### 454 **Data availability**

455 The datasets used in the current study are available from DJFR; [dr60@st-andrews.ac.uk](mailto:dr60@st-andrews.ac.uk).

456

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641 **Author contributions statement**

642 Conceived the study: MIDC, DJFR, CBE, KAB. Wrote the manuscript: MIDC. Contributed data: DT, KAB.  
643 Processed the data: MIDC, DJFR, CJB. Analysed the data: MIDC. Advised data analysis: DJFR, CBE, PJH. All  
644 authors reviewed and edited the manuscript.

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646 **Additional information**

647 The authors declare no competing interests, financial or otherwise.

648 **Tables**

649 **Table 1: Device deployment summary information.** Tagged pup sample sizes and tag duration by  
 650 deployment site and year. Trip and dive numbers given are those included in the analysis after data  
 651 cleaning and restriction to 120 days after leaving the colony. Although SRDL devices recorded dives, these  
 652 could not be matched to bathymetric depth data and so were excluded from dive analysis. Colonies were  
 653 assigned to two geographic regions; NS = North Sea, CIS = Celtic and Irish Seas.

Deployment site (year)	Region	Device type	No. tagged seals			Mean no. locations day <sup>-1</sup> ± SD	Total no. trips	Total no. dives
			f	m	Total			
Isle of May (2001)	NS	SRDL	5	6	11	4.5 ± 2.3	109	N/A
Isle of May (2002)	NS	SRDL	5	5	10	5.2 ± 1.7	67	N/A
Bardsey (2009)	CIS	GPS-GSM	2	0	2	35.5 ± 5.4	23	3871
The Skerries (2009)	CIS	GPS-GSM	1	2	3	33.1 ± 5.7	141	9373
The Skerries (2010)	CIS	GPS-GSM	4	1	5	57.2 ± 13.8	212	46589
Ramsey (2010)	CIS	GPS-GSM	3	4	7	37.3 ± 9.7	162	27609
Muckle Green Holm (2010)	NS	GPS-GSM	4	3	7	22.5 ± 9.6	38	7417
Stroma (2010)	NS	GPS-GSM	5	2	7	24.4 ± 4	84	7941
		<b>Total:</b>	<b>29</b>	<b>23</b>	<b>52</b>		<b>836</b>	<b>102800</b>

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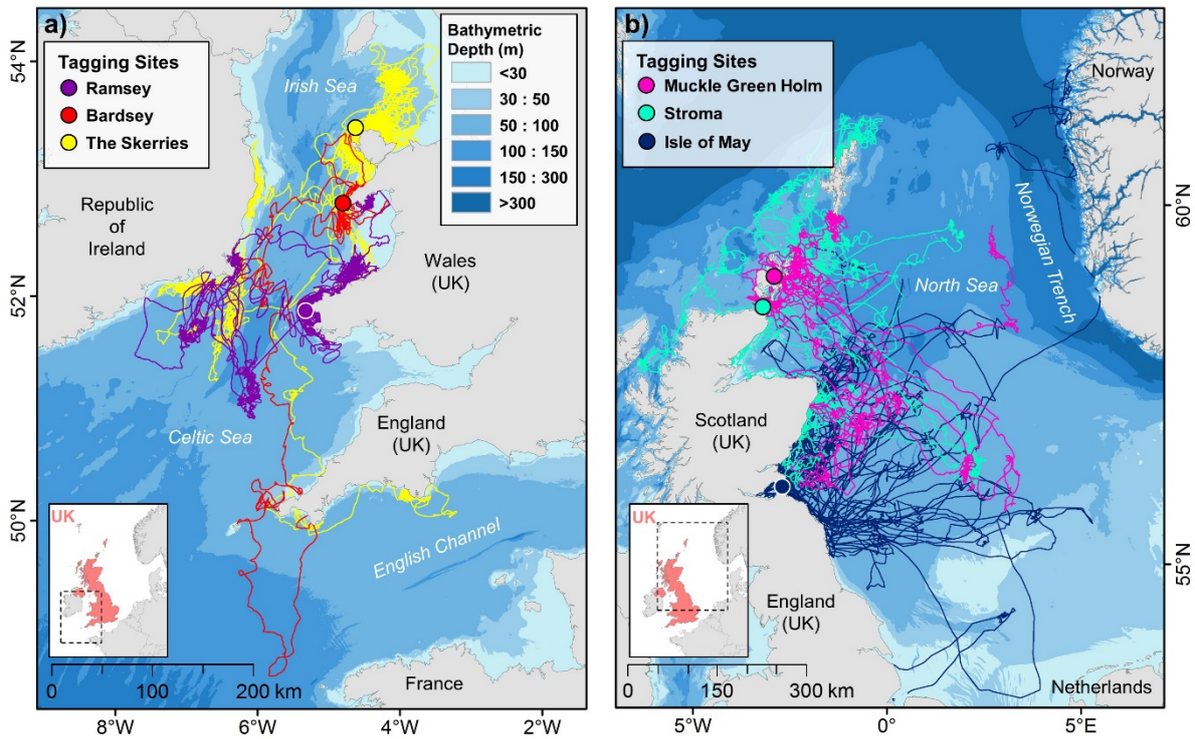
Accepted

655 **Table 2: Model output.** Results of model simplification using backwards hypothesis testing with GEE-GAMs. Significant ( $p < 0.05$ ) terms are shown with “\*”.  
 656 Interactions between variables are denoted by “:”. Where a variable was significant in an interaction, the significance of component interactions and/or individual  
 657 fixed effects is not reported.

Predictor Variables	Response Variables							
	Trips		Dives (daily means)					
	Duration (Fig. 4a-b)	Distance (Fig. 4c-d)	Max. Depth (Fig. 5a-b)	Bathy. Depth (Fig. 5c-d)	Prop. Benthic (Fig. 5e-f)	Duration (Fig. 6a-b)	Prop. Bottom Time (Fig. 6c-d)	Prop. Time Diving (Fig. 6e-f)
<b>Time</b>	$\chi^2_3 = 15.2, p = 0.002^*$	$\chi^2_3 = 8.2, p = 0.042^*$	-	-	-	-	-	-
<b>Sex</b>	$\chi^2_1 = 1.4, p = 0.233$	-	-	-	$\chi^2_1 = 5.2, p = 0.023^*$	$\chi^2_1 = 2.5, p = 0.117$	-	-
<b>Region</b>	$\chi^2_1 = 66.1, p < 0.001^*$	-	-	-	-	-	-	-
<b>Time : Sex</b>	$\chi^2_3 = 3.9, p = 0.268$	$\chi^2_3 = 6.2, p = 0.1$	-	-	$\chi^2_3 = 1.1, p = 0.774$	$\chi^2_3 = 3.1, p = 0.369$	$\chi^2_3 = 3.1, p = 0.378$	$\chi^2_3 = 13.9, p = 0.003^*$
<b>Time : Region</b>	$\chi^2_3 = 4.1, p = 0.254$	$\chi^2_3 = 3.3, p = 0.346$	-	-	$\chi^2_3 = 13.1, p = 0.004^*$	$\chi^2_3 = 16.4, p < 0.001^*$	$\chi^2_3 = 14.9, p = 0.002^*$	$\chi^2_3 = 15, p = 0.002^*$
<b>Region : Sex</b>	$\chi^2_1 = 2.7, p = 0.099$	$\chi^2_1 = 4.7, p = 0.03^*$	-	-	$\chi^2_1 = 0.1, p = 0.767$	$\chi^2_1 = 0, p = 0.875$	$\chi^2_1 = 9.3, p = 0.002^*$	$\chi^2_1 = 0.02, p = 0.885$
<b>Time : Region : Sex</b>	$\chi^2_3 = 0.8, p = 0.852$	$\chi^2_3 = 1.4, p = 0.708$	$\chi^2_3 = 13.6, p = 0.003^*$	$\chi^2_3 = 10.4, p = 0.016^*$	$\chi^2_3 = 1.3, p = 0.74$	$\chi^2_3 = 1.9, p = 0.591$	$\chi^2_3 = 7.24, p = 0.065$	$\chi^2_3 = 4.5, p = 0.215$

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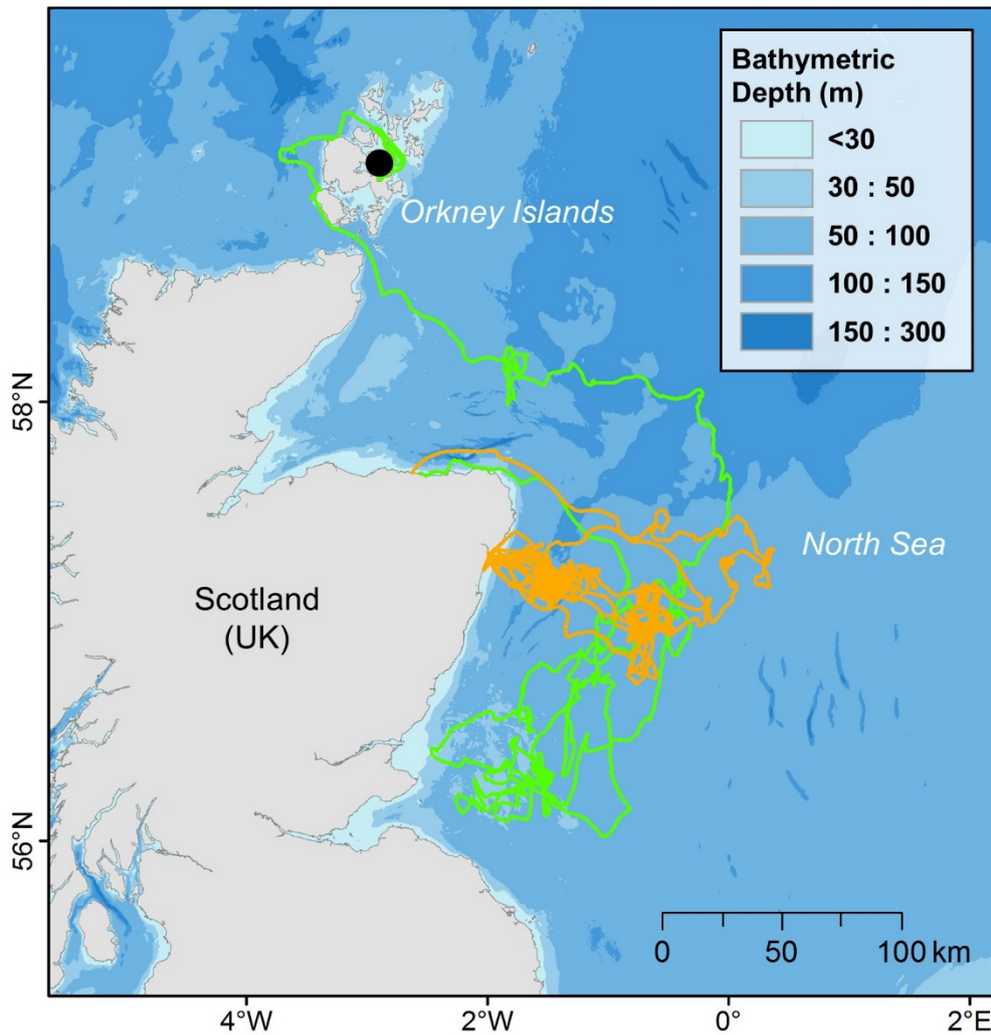
659 **Figure Captions**



660

661 **Figure 1: Pup tagging sites and trips at sea.** Pups were tagged at six colonies in the United Kingdom (UK).  
662 Colonies were assigned to one of two geographic regions; (a) Celtic and Irish Seas (CIS), and (b) North Sea  
663 (NS). Tracks show pup trips (n=836) during the initial four months after leaving the colony. Maps created  
664 in Esri ArcMap™ 10.2.2 (<http://desktop.arcgis.com/en/arcmap/>).

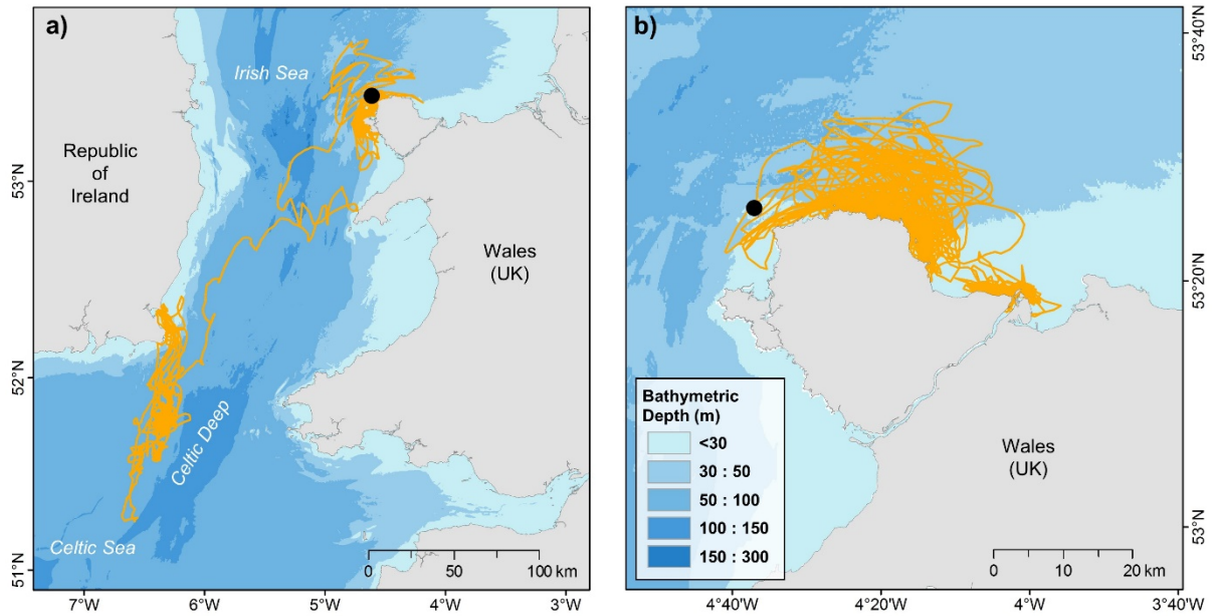
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667 **Figure 2: Changes in NS pup trip behaviour with time since departing the colony.** Map shows initial  
 668 exploratory trip of a pup from Muckle Green Holm, Orkney Isles (black dot), during which it did not haul-  
 669 out for 64 days (green track). During the following 56 days, the pup performed repeated short-duration (5-  
 670 14 days) foraging trips (gold tracks), travelling between the haul-out site and specific putative foraging  
 671 areas. Map created in Esri ArcMap™ 10.2.2 (<http://desktop.arcgis.com/en/arcmap/>).

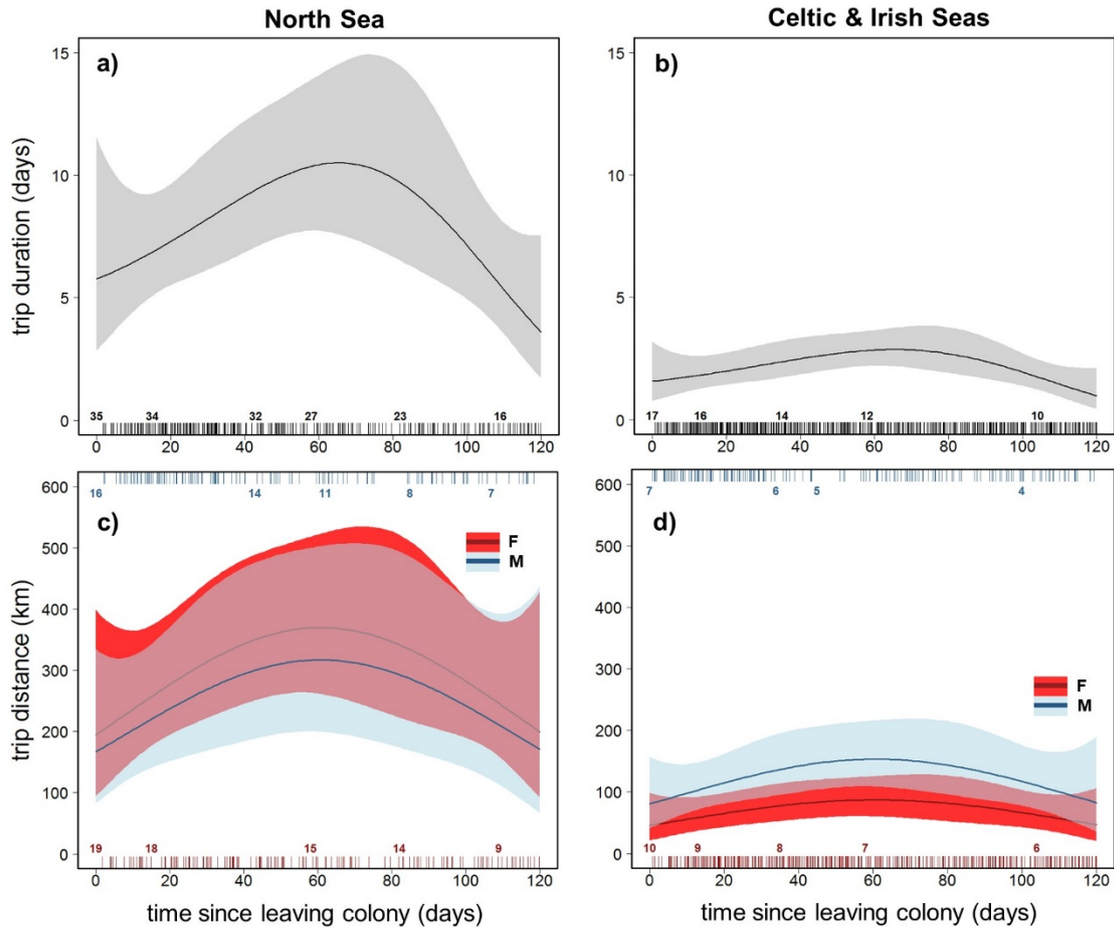
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674 **Figure 3: Differences in dispersal of CIS pups.** Maps show trips of two pups from The Skerries over the  
 675 first 4 months of independence. Pup (a) remains in areas adjacent to the colony (black dot) for ~50 days  
 676 before dispersing to the southeast coast of Ireland, hauling-out at a popular grey seal haul-out site, and  
 677 subsequently making repeated trips to putative foraging grounds on the edge of the Celtic Deep. Pup (b)  
 678 remains in areas adjacent to the colony (black dot) for the entire 4 months. Maps created in Esri ArcMap™  
 679 10.2.2 (<http://desktop.arcgis.com/en/arcmap/>).

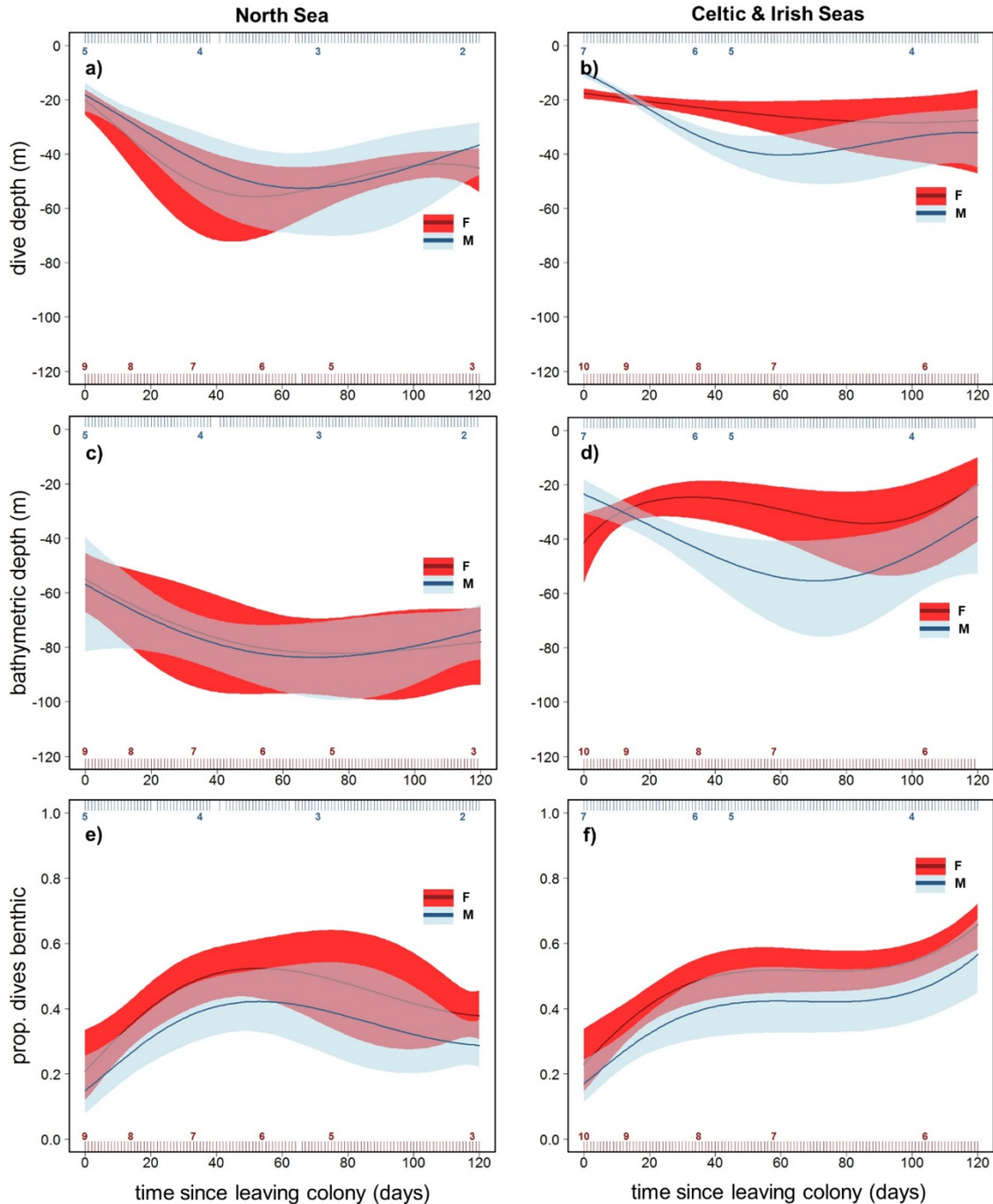
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682 **Figure 4: Sex and region differences in ontogeny of trip behaviour.** Model-fitted values for trip  
 683 duration (a-b) and trip distance (c-d) over time since leaving the colony. Solid lines show population mean  
 684 responses by region (North Sea (NS) left, Celtic and Irish Seas (CIS) right), with associated GEE-based 95%  
 685 confidence intervals (shaded areas). Pup trip behaviour changed significantly with time since departure.  
 686 NS pups performed longer duration trips than CIS pups, however there was no sex difference (a-b). CIS  
 687 males (d; blue) travelled further than females (red). Rug plots top and bottom show the distribution of data,  
 688 colour-coded by sex, and associated numbers indicate pup sample size.

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690

691 **Figure 5: Sex and region differences in ontogeny of dive depth.** Model-fitted values for daily mean  
 692 maximum dive depth (a-b) and bathymetric depth of dive locations (c-d) over time since leaving the colony.  
 693 Solid lines show population mean responses by region (North Sea (NS) left, Celtic and Irish Seas (CIS) right),  
 694 with associated GEE-based 95% confidence intervals (shaded areas). Pups increased their dive depth  
 695 rapidly over the initial 40 days (a-b), except for CIS females (b). NS pups dived in deeper water throughout  
 696 (c). Sex differences in bathymetric depth of dive locations emerged from the outset in CIS pups, as females  
 697 (red) dived in shallower areas (d). The proportion of dives that were benthic increased rapidly for all pups  
 698 over the initial 40 days. However, females recorded marginally higher mean values than males in both

699 regions (e-f). Rug plots top and bottom show the distribution of data, colour-coded by sex, and associated  
700 numbers indicate pup sample size.

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