
Ecological bridges and barriers in pelagic ecosystems

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Abstract :

Many highly mobile species are known to use persistent pathways or corridors to move between habitat patches in which conditions are favorable for particular activities, such as breeding or foraging. In the marine realm, environmental variability can lead to the development of temporary periods of anomalous oceanographic conditions that can connect individuals to areas of habitat outside a population's usual range, or alternatively, restrict individuals from areas usually within their range, thus acting as ecological bridges or ecological barriers. These temporary features can result in novel or irregular trophic interactions and changes in population spatial dynamics, and, therefore, may have significant implications for management of marine ecosystems. Here, we provide evidence of ecological bridges and barriers in different ocean regions, drawing upon five case studies in which particular oceanographic conditions have facilitated or restricted the movements of individuals from highly migratory species. We discuss the potential population-level significance of ecological bridges and barriers, with respect to the life history characteristics of different species, and inter- and intra-population variability in habitat use. Finally, we summarize the persistence of bridge dynamics with time, our ability to monitor bridges and barriers in a changing climate, and implications for forecasting future climate-mediated ecosystem change.

Keywords : Species distribution, Migration corridors, Population connectivity, Oceanographic features, Tuna, Billfish, Marine mammal, Brazilian episode

56

1. Introduction

58 Throughout the biosphere and across all scales of ecological organization, the
60 environmental conditions that constitute animal habitats are arranged in a complex,
62 hierarchical and heterogeneous configuration. This patchiness can restrict sessile animals
64 to the same habitat 'patch' for most of their lifetime while mobile animals can move between
66 suitable patches, because they can tolerate unfavorable conditions when moving between
68 preferred habitats (Switzer, 1993). These movements across a landscape or seascape
70 connect populations and allow for life history processes that are essential to population
72 persistence (Gilbert-Norton et al., 2010) and ecosystem function (e.g. genetic flow, nutrient
74 cycling, Reimchen et al. (2003); Sanchez-Pinero and Polis (2000)).

68 Habitat connectivity – the degree of interconnectivity between patches of favorable
70 habitat – is essential, not only for individual survival but also for the maintenance of
72 metapopulation structure, and ultimately, biological diversity (Hanski, 1999). Seasonal
74 events may trigger dispersal or migration to exploit different habitats that are beneficial to
76 reproduction and fitness (Dingle, 2009; Murrell et al., 2002). In terrestrial landscapes, many
78 large-bodied ungulates and winged species undergo lengthy migrations (Harris et al.,
80 2009), avoiding unsuitable habitats, crossing barriers or temporarily tolerating unsuitable
82 environments (e.g. wildebeest in Serengeti (Ottichilo et al., 2001) and raptor migrations
84 across Sahara (Strandberg et al., 2009)).

78 In marine systems, satellite-tracking studies of pelagic fishes, sea turtles, seabirds
80 and marine mammals have shown impressive transoceanic migrations between areas used
82 for different stages of the ontogenetic or annual cycle (Akesson and Hedenstrom, 2007;
84 Block et al., 2011; Bonfil et al., 2005; Scott and Hays, 2014; Shaffer et al., 2006). The
86 routes that characterize movement between suitable habitats and that are spatially
88 persistent are known as corridors (Anderson et al., 2013; Beier and Noss, 1998; Bennett,
90 1999). While habitat corridors in terrestrial environments are well understood (e.g. monarch
92 butterflies (Brower, 1995); osprey (Alerstam et al., 2006)), the concept of corridors in the
94 marine realm is less developed. Yet highly mobile marine species are also known to utilize
96 seasonally dynamic oceanographic features to move between known breeding and foraging
98 habitats (Guilford et al., 2009; Morreale et al., 1996; Polovina et al., 2006). Some corridors
100 are well defined by the seasonal or annual predictability of a population returning
102 generation after generation (Anderson et al., 2013).

92 There are also locations within a species range that have periodic bouts of
94 anomalous environmental conditions that may influence habitat suitability. In pelagic
96 systems, currents and mesoscale oceanographic features (e.g. eddies, fronts, filaments,
98 changes in vertical mixing) are the major sources of this environmental variability over intra-
100 annual timescales (Bakun, 2006). In contrast to predictable and regularly used migratory
102 corridors, anomalous environmental conditions may lead to the development of short-lived
104 corridors or *ecological bridges*. Following Fromentin et al. (2014a), we define an ecological
106 bridge as a temporary habitat pathway connecting two suitable but distinct habitat regions
108 (Fig. 1). Anomalous oceanographic conditions and changes in mesoscale variability can
110 create such ecological bridges, and allow individuals access to alternate, or irregular, areas
112 of habitat. We distinguish this from cases where a single habitat expands to new regions
114 thereby allowing species to increase their range (e.g. Mackenzie et al., 2014; Stewart et al.,

104 2014), which has been increasingly observed as ecosystems respond to global climate
change (Hollowed et al., 2013; Kirby et al., 2006).

106

In contrast to ecological bridges, migrating animals are often confronted with
108 barriers between favorable habitat patches. Ecological barriers can be geographic (e.g.
seas, land masses, deserts, or mountains), or environmental (e.g. temperature and salinity
110 gradients, light or oxygen levels, (Prince and Goodyear, 2006; Selkoe et al., 2008).
Profound changes to corridors and barriers have occurred in the past, e.g. historical
112 episodes of climate change and tectonic activity (Gaston, 2003). Some change more
quickly, in synchrony with timing and intensity of interannual and decadal events (ENSO
114 and PDO) (Lehodey et al., 1997; Massom and Stammerjohn, 2010).

116 The timing and location of ecological bridges and barriers may change over space
and time, connecting (or disconnecting) animals to disjunct (or adjunct) habitats, with a
118 range of ecological implications. Here we provide evidence of ecological bridges and
barriers in the marine realm, drawing upon case studies in which a particular set of
120 oceanographic conditions have facilitated or prevented the movements of individuals
between patches of favourable habitat. In Section 2, we present five case studies of
122 ecological bridges and barriers, detailing how each bridge (barrier) is formed and how
species respond, how the presence of a bridge (barrier) affects population structure and
124 connectivity, and the socio-economic implications (if any). We then generalise the
importance of bridges and barriers in terms of dynamics, population level significance, and
126 future research needs (Section 3).

128 **2. Ecological bridges and barriers in pelagic systems**

The movements and migratory patterns of pelagic species can have important
130 ecological and population level effects (Dingle, 2014; Frisk et al., 2014), especially in higher
trophic level predators which can play an important role in structuring and maintaining
132 marine food webs (Heithaus et al., 2008). While both physical and biological factors
influence the movements and resulting patterns in population structure and connectivity in
134 marine species (Frisk et al., 2014), we mostly focus here on changes in the physical
environment. Case studies from pelagic fishes and marine mammals demonstrate how
136 shifting environmental conditions create ecological bridges or barriers that can influence the
distribution of migratory marine species with potentially important ecological effects at the
138 population level, as described below.

140 **2.1 Atlantic bluefin tuna: the Brazilian episode and a bridge between two hemispheres**

142 Throughout its thousand years of exploitation, catches of Atlantic bluefin tuna
(ABFT, *Thunnus thynnus*) have exhibited conspicuous changes in both time and space
144 domains (Fromentin et al., 2014a; Mather et al., 1995; Ravier and Fromentin, 2004),
probably reflecting the high mobility of the species (Block et al., 2005; Sibert et al., 2006).
146 During the 20th century, large Nordic and Japanese fisheries rapidly arose in unexpected
fishing areas, i.e. the North and Norwegian Seas and the equatorial Atlantic, but suddenly
148 disappeared after a few years or decades. Those variations seem to be primarily due to
environmentally driven changes in ABFT migration patterns that could act in synergy with
150 local/regional overfishing (Fromentin, 2009).

152 One of the most striking changes in ABFT spatial distribution was the so-called
154 “Brazilian episode”, during which Japanese longline fishing boats caught large quantities of
156 ABFT (a temperate species) in the equatorial Atlantic where they were targeting tropical
158 tunas (Fromentin et al., 2014a; Takeuchi et al., 2009). In a study by Fromentin et al.
160 (2014a), a niche model was applied to an extensive dataset of catch and environmental
162 variables from 1960 to 2009. Results showed that ABFT has a remarkably large ecological
164 niche, with high probabilities of occurrence in the North Atlantic and adjacent seas (as
166 expected), as well as in the South Atlantic at around 30°S and along the southwestern
168 African coast (Fig. 2a). The niche model also detected favorable environmental conditions
170 for ABFT in the western equatorial Atlantic during the 1960s, exactly where the Japanese
172 vessels caught ABFT. The 1960s were the only decade in the last 50 years that exhibited
relatively high probability of ABFT occurrence around the Equator. No ABFT have been
caught in the equatorial Atlantic since then, although the fishing effort significantly increased
in that area. During the last decade, higher probabilities of ABFT mostly occurred above
45°N (Fig. 2b), which could be related to global warming and which agrees with a northward
expansion of ABFT (see below). ABFT could have thus migrated from their northern
spawning grounds to the South Atlantic during the 1960s through the western equatorial
Atlantic acting as an ecological bridge between the central North and the central South
Atlantic. These new geographical spots could have subsequently been transmitted from
year-to-year through spatial learning and entrainment of younger fish (Petitgas et al., 2010).

Further analyses indicated that during that period, ABFT could have migrated from
the equatorial Atlantic to the western spawning ground of the Gulf of Mexico during the first
part of the year followed by a reverse north-south migration during the second part of the
year (Fromentin et al., 2014a). The southeastern Atlantic feeding grounds (offshore of
South Africa, Namibia and Angola) may well have been shared by both ABFT and southern
bluefin tuna during the 1960s. However, this bridge appears to have broken by the late
1960s because of oceanographic changes affecting primarily sea surface temperature and
possibly the equatorial current and counter-current. This could have made ABFT migration
to the South Atlantic more difficult. ABFT has high rates of natal homing (Rooker et al.,
2008) and as connections have been detected between the western equatorial Atlantic and
the Gulf of Mexico, but not with the Mediterranean Sea, it is highly probable that those
individuals were part of the western Atlantic ABFT stock. Therefore, the breaking of this
ecological bridge may have primarily affected the productivity of the western stock and its
lack of rebuilding could result from a regime shift due to the combination of oceanographic
changes in the equatorial Atlantic (the breaking of the ecological bridge) and intense fishing
in the North Atlantic in the 1960s-1970s (Fromentin et al., 2014a).

190 Because ABFT displays a large ecological niche, it has potentially more abiotic
192 opportunities (i.e. a larger ecological window) than many other large pelagic fish. This may
194 explain why the ABFT spatial distribution appears generally highly variable. The ABFT
196 spatial distribution seems to have expanded northward (beyond 50°N) in the last decade
198 (as it did from the 1930s to the 1950s), probably because of the effects of global climate
change. This is particularly evident in the western Atlantic, with the northern expansion of
large ABFT in the Gulf of Saint Lawrence. MacKenzie et al. (2014) postulated that the
presence of bluefin tuna in waters east of Greenland in 2012 could be due to a combination
of warmer temperatures and immigration of an important prey species (mackerel) to the

200 region, indicating that global warming will open (or close) ecological bridges in the northern
(southern) parts of the oceans to marine fish and, probably to other marine vertebrates.

202 **2.2 Atlantic bluefin tuna: the western pulse into the Bay of Biscay**

204 In the Northeast Atlantic, the Bay of Biscay is a key juvenile ABFT feeding ground.
206 Juveniles migrate into the Bay of Biscay in April-June, where they feed on the abundant
208 local prey and grow before migrating back to Atlantic wintering areas in autumn (Dufour et
210 al., 2010). Some adults also feed in the Bay of Biscay after spawning in the Mediterranean
(Aranda et al., 2013). Using different technologies (such as conventional tags, electronic
212 tags, or chemical tags), Bay of Biscay ABFT have been shown to be substantially
214 connected with other areas across the Mediterranean and the western Atlantic (Abascal et
216 al., 2016; Arregui et al., 2006; Graves et al., 2015). However, the natal origin of Bay of
218 Biscay ABFT remained unknown until the otolith chemistry study by Fraile et al. (2015).
220 Using a substantial sample of juveniles and adults over three consecutive years, they found
222 that the Bay of Biscay is supported almost exclusively by the eastern ABFT population, but
the western population may also occasionally contribute in some years. Given that the
eastern population is believed to be an order of magnitude larger than the western
population (Kerr et al., 2015), ABFT of western origin is particularly difficult to detect in
eastern foraging areas. In fact, a small proportion of western migrants was detected in the
Bay of Biscay only in 2009 – a western pulse. In contrast, the Bay of Biscay fishery was
composed exclusively of eastern origin bluefin tuna in 2010 and 2011. Based on their
sampling, Fraile et al. (2015) suggested that a substantial fraction of the western population
may move across the Atlantic Ocean to feed in the Bay of Biscay and/or surrounding waters
of the Northeast Atlantic.

224 Across the three consecutive fishing seasons, all the western origin ABFT detected
226 in the Bay of Biscay were caught within a very restricted time window (10 days) in 2009,
228 suggesting high temporal variability in the transatlantic migration from west to east, with
migration events occurring in sporadic pulses that could be related to variability in
230 environmental conditions (Fraile et al., 2015). A recently developed habitat model that
232 notably includes productive mesoscale features as a proxy for food availability (Druon et al.,
234 2016) suggests that the 2009 pulse of western origin ABFT into the Bay of Biscay might
236 have been due to the existence of a longitudinal ecological bridge across the Atlantic (Fig.
238 3a). One to three months prior to sampling in 2009, this habitat bridge which is linked to the
240 Gulf Stream dynamics connected the main western and eastern Atlantic feeding areas
through a well-defined, relatively narrow corridor west of 45°W. During the ABFT migration
period to northeast Atlantic feeding grounds after wintering (from April to June), the
potential feeding habitat in the central part of the bridge was observed to be largest in 2009
compared to 2010 and 2011 (Fig. 3d). The bridge between the eastern and western feeding
areas in 2010 and 2011 was less marked and more discontinuous, which might have acted
as a barrier against migration of western origin ABFT into the eastern Atlantic feeding
grounds (Fig. 3b and c).

242 Mixing of eastern and western ABFT across the whole Atlantic Ocean remains one
244 of the most critical uncertainties preventing accurate diagnoses of stock status to guide
246 effective management (Fromentin et al., 2014b). Different studies have illustrated the
complexity of the connectivity between remote Atlantic areas and their implications for
ABFT management (Block et al., 2005; Galuardi and Lutcavage, 2012; Rooker et al., 2014;

248 Rooker et al., 2008). Effective fishery management will require a better understanding of the
249 magnitude of these movements, their temporal variability, and the physical and biological
250 factors that may affect it (Graves et al., 2015).

250

251 More research is needed to better understand the role of habitat bridges and
252 barriers in relation to ABFT population connectivity. If bridges are not persistent over time
253 as driven by climate change over the Gulf Stream dynamics, western origin ABFT that
254 migrated into the eastern Atlantic might be less likely to return to the west (and vice versa
255 for the eastern origin population to the east). Depending on the magnitude of such potential
256 habitat barriers preventing the migration back to the west, important implications could
257 include exposure to potentially higher fishing intensity, or delays in the natal homing
258 behavior to spawn in the Gulf of Mexico (and vice versa in the Mediterranean Sea for the
259 eastern origin population). The monitoring of these habitat contraction and relaxation acting
260 as barriers and bridges under climate change is therefore essential to evaluate the
261 important potential implications for ABFT population connectivity and dynamics.

262

263 **2.3 Southern bluefin tuna: pathways to southeast Australia**

264 Migration pathways of southern bluefin tuna (SBT *Thunnus maccoyii*) have been
265 studied over many decades with conventional, acoustic, archival and satellite tagging
266 programs, providing a range of insights into their movement and behaviour (Bestley et al.,
267 2009; Fujioka et al., 2010; Hobday et al., 2015; Patterson et al., 2008). After hatching, fish
268 move from the single-known spawning ground between Indonesia and Australia, following
269 the Leeuwin Current down the Australian west coast to reach the southern coast by age 1.
270 They are then resident during the austral summers in the Great Australia Bight (GAB)
271 between the ages of 2-5 years. At the end of each summer, juvenile SBT leave the GAB
272 and move east to the Tasman Sea or west to the Indian Ocean where they spend the winter
273 feeding (Bestley et al., 2009).

274 A possible ecological bridge connecting juvenile SBT habitats was interrupted in the
275 early 2000s, when eastward migration to the Tasman Sea became rare. Conventional tag-
276 recapture data revealed that fewer juvenile SBT tagged in the 2000s moved into the
277 Tasman Sea compared to fish tagged in the 1990s (Basson et al., 2012). Based on
278 thousands of tag returns from SBT tagged at ages 1 and 2, the percent of returns coming
279 from the Tasman Sea was much higher in the 1990s (5.7% and 12.8% for age 1 and 2
280 respectively) than in the 2000s (1% and 0.4% for age 1 and 2, respectively). Archival tag
281 tracks also provide evidence for reduced eastward movement of juvenile SBT in the 2000s
282 (Basson et al., 2012). Only 4% of tracks (3 out of 75) showed movement into the Tasman
283 Sea (>150°E) during the months of May through November after 2001, compared to 21%
284 (14 out of 67) in prior years (chi-squared test p-value=0.01). The exact timing of this change
285 is difficult to determine as few tags returned data between 2001 and 2004.

286 These migration pathway changes may be in response to population decline (there
287 has been a documented decline in SBT abundance and recruitment through the 1990s and
288 into the 2000s, and cohorts in 2000-2002 were at historically low levels (Anon, 2009)), or to
289 environmental changes that affect SBT migration. In the Tasman Sea, a long-term warming
290 trend has been observed (Hobday and Pecl, 2014; Ridgway, 2007). Other areas occupied
291 by juvenile SBT, such as the GAB and eastern Indian Ocean have not warmed as rapidly
292 over the same period (Basson et al., 2012; Hobday and Pecl, 2014). This warming may be
293 acting as a partial barrier to restrict juvenile SBT movements to areas they occupied in the
294 1990s.

296 Habitat models for juvenile SBT have been developed using location data collected
298 on SBT over many years from electronic tags, and comparing the ocean conditions where
300 fish were found with the conditions available to them throughout the region and time period
302 of interest (Basson et al., 2012; Eveson et al., 2014). Sea surface temperature (SST) and
304 chlorophyll were found to have the greatest influence. Habitat models based on SBT
306 preferences for SST and chlorophyll revealed a high preference habitat band in April to
308 June along the west and south coasts of Tasmania into the Tasman Sea in the period 1998-
310 2000 that was no longer present in the period 2004-2006 (Basson et al., 2012). The habitat
312 models have subsequently been updated to include new archival tag data and to use
314 improved fish location estimates based on a recently published method for light-based
316 geolocation (Basson et al., 2016). These updated models continue to show the
318 disappearance of a connecting habitat band between the GAB and the Tasman Sea
between the two time periods (Fig. 4). We note that separate habitat models were used for
the two time periods since habitat preferences for SBT changed slightly between these
periods (Fig. 4a,b). If we use a single habitat model based on the entire period 1998-2006,
the missing band of suitable habitat in 2004-2006 is still evident but less extreme. We argue
it is more defensible to use separate habitat models than to combine all years into a single
model that masks the preference change. This does, however, raise the dilemma of
whether the observed ecological barrier has arisen due to changes in fish physiology and
behaviour or to environmental changes – most likely a combination of both. Changes in
additional variables, such as forage distribution, remain difficult to estimate, and the habitat
model remains a proxy for environmental change that restricted movements of juvenile SBT
across this ecological bridge.

318 This example of an ecological bridge “breaking” foreshadows changes that are
320 expected under climate change as environmental tolerances are exceeded in some regions
322 (Burrows et al., 2014). If the ecological barrier persists, the implications for SBT populations
324 are likely to be relatively minor, as this region is only a small part of their total range and
326 larger SBT may not be restricted by the barrier. Dependent fisheries in eastern Australia
and New Zealand may experience declines in catch, however, we are unable to estimate
these effects. A new archival tagging program in the GAB commenced in 2015, and in a few
years will allow new estimates of east-west migration and assessment of the state of this
ecological bridge.

328 **2.4 Blue marlin: intermittent crossing of the Equatorial Pacific**

330 Like bluefin tuna, blue marlin (*Makaira nigricans*) is a wide-ranging species, with
332 some of the most impressive long-range movements ever recorded for oceanic fishes
334 (Kraus et al., 2011; Ortiz et al., 2003). In the North Pacific, 59 marlin were tracked from
2009-2013 moving south from Hawaii, crossing the equator and moving towards French
Polynesia (Fig. 5). For most migratory species, the equator serves as a natural ecological
barrier (e.g. see ABFT example above), due to the combination of high sea-surface
temperature and oxygen limits at relatively shallow depths (MacLeod, 2009). However, in a
recent study blue marlin were shown to routinely undergo a unique, trans-equatorial
migratory strategy (Carlisle et al., In Press).

338 Interestingly, this trans-equatorial route was not used by blue marlin in 2010,
340 perhaps due to a La Nina event. This cold phase of the ENSO cycle, which in the North
Pacific is characterized by a western extension of the cool SST water mass from the
342 eastern Pacific (cool tongue), increased equatorial upwelling and shoaling of the
thermocline and oxycline (Philander, 1989; Wyrki, 1975). Blue marlin tagged in 2010

344 moved south as they did during other years until they encountered the western extension of
the cool tongue (Fig. 5b), which had water temperatures below 24°C, below their preferred
346 thermal range of 26 to 30°C (Goodyear et al., 2008; Graves et al., 2001; Holland et al.,
1990). Upon encountering the cool tongue, the blue marlin stopped moving south and
348 remained in the warm waters to the north of this cold oceanographic feature, with several
fish moving longitudinally along its northern boundary. These cold temperatures, combined
350 with the increased vertical habitat compression associated with shoaling of cold, low oxygen
waters driven by increased equatorial upwelling, appeared to present a vertical and
horizontal ecological barrier to trans-equatorial movements. During non-La Niña years, this
352 oceanographic barrier to trans-equatorial migration is not present as SSTs are not limiting
(Fig. 5a) and vertical habitat compression is reduced.

354 The effect on the population dynamics of blue marlin will depend on the extent and
persistence of the barrier as well as the nature of the trans-equatorial migration. The
356 purpose of the trans-equatorial migrations of blue marlin remains unclear, but in general the
broad-scale migratory patterns of blue marlin have been linked to foraging and reproductive
358 migrations (Shimose et al., 2009; Shimose et al., 2012). Hawaii is a known spawning
location (Hopper, 1990; Seki et al., 2002), and French Polynesia has also been identified as
360 a region where spawning occurs (Howard and Ueyangi, 1965). Hence, trans-equatorial
movements may be related to spawning and disruption of these potential spawning
362 migrations may have important effects in terms of population connectivity. In addition,
reduced mixing rates between different populations may increase susceptibility of blue
364 marlin to localized depletion due to overfishing (Lee et al., 2014). Much remains unclear
about how oceanographic conditions will be altered under future climate change, but some
366 research suggests that there will be an increase in the frequency of extreme El Niño and La
Niña events (Cai et al., 2014; Cai et al., 2015; Power et al., 2013). Any increase in the
368 intensity or frequency of La Niña events will likely increase the extent and persistence of
such barriers, potentially dividing the population of blue marlin in the Central Pacific.

370

2.5 Bowhead whales: traversing the Northwest Passage

372 The Northwest Passage (NWP) is a series of Arctic waterways connecting the
Atlantic and Pacific Oceans. Throughout most of the year, dense ice cover within the NWP
374 represents a physical barrier between the two oceans (McKeon et al., 2015). Arctic species
are well adapted to such barriers, and have tuned their feeding and breeding behaviors to
376 coincide with seasonal changes in ice pack. Bowhead whales (*Balaena mysticetus*) are the
largest Arctic predator, with a wide-ranging distribution and populations found on both sides
378 of the NWP. The species is well suited for ice-covered waters, given their ability to move
through extensive areas of sea ice coverage (Citta et al., 2015; George et al., 1989; Heide-
380 Jørgensen et al., 2012; Laidre et al., 2008). Individuals spend the summer months foraging
in Arctic waters and then migrate to subarctic seas during the winter months (Laidre et al.,
382 2008).

While genetic evidence indicates historic gene flow between Atlantic and Pacific
384 populations (Alter et al., 2012), the lack of bowhead remains from interior locations in the
NWP suggests that individuals have maintained separate populations (McKeon et al.,
386 2015). However, in the summer of 2010, and following a long-term warming trend, the NWP
was suitably free of ice to allow two individuals from separate populations to forage in the
388 same region at the same time (Heide-Jørgensen et al., 2012; McKeon et al., 2015).
Individuals migrated back to their respective oceans after ten days. However, this short

390 occupation of common territory demonstrated the occurrence of an ecological bridge,
through which bowhead whales were capable of inter-population exchange based on sea-
392 ice conditions (Heide-Jørgensen et al., 2012; McKeon et al., 2015).

The extent and thickness of Arctic sea ice has continued to decrease at an alarming
394 rate (McKeon et al., 2015), and the accelerated loss of sea ice will increase the ease and
frequency with which marine species are able to move between the Pacific and Atlantic
396 Ocean basins (Heide-Jørgensen et al., 2012). As such, the disappearance of long standing
ice barriers and subsequent increased frequency of bridge conditions will have a dramatic
398 impact on a range of Arctic species (McKeon et al. 2015). As the effects of climate-
mediated ecosystem change are likely to be most pronounced in the Arctic in upcoming
400 decades (Burrows et al., 2014; Moore and Huntington, 2008), it is perhaps not surprising
that bridges and barriers will appear in this region. The dynamics of ice-melt and the effects
402 on availability of preferred foraging habitats will see Arctic marine mammals and seabirds
begin to explore novel areas (McKeon et al., 2015). At the same time, greater accessibility
404 to humans (e.g. increased ship transport, oil exploration, and industrial fishing) may have
serious ecological impacts for Arctic species (McKeon et al., 2015).

406

3. Importance of ecological bridges and barriers

408 The preceding examples illustrate that highly migratory pelagic species encounter
ecological bridges and barriers that have facilitated or prevented individual movements over
410 a range of space and time scales. In pelagic systems, the range over which individuals from
a population tend to roam is an important consideration in the ability of those individuals to
412 exploit an ecological bridge, or be restricted by an ecological barrier. The case studies
presented here describe the movements of large teleost fish and marine mammals, which
414 are among the most wide-ranging of all pelagic marine vertebrates (Block et al., 2011) and
so most likely to encounter novel habitat conditions. Ecological bridges and barriers can
416 modify spatial dynamics and connectivity of a population, impact on fisheries, and in the
long term may affect population structure. For example, connectivity to new habitat may
418 initiate conspecific interactions between separate populations, introduce new competition
for resources, and modify existing biotic interactions and phenotypic traits (Brown et al.,
420 2015). Below, we discuss the persistence in bridge dynamics with time, individual to
population level sensitivity, and our ability to monitor bridges and barriers in a changing
422 climate.

3.1 Bridge and barrier dynamics

424 Importantly, ecological bridges and barriers may support a complex meta-population
426 structure and thus safeguard populations from local extinction events (e.g. hypoxic dead
zones, corrosive waters), inter-annual variability (e.g. ENSO-related events, 'anomalous'
428 years), and even unprecedented changes to oceans. As rapid climate change is expected
to impact pelagic species (Dell et al., 2015; McBride et al., 2014; Robinson et al., 2015), the
430 spatio-temporal dynamics of ecological bridges and barriers will be inherently linked to the
periodicity and frequency of environmental and oceanographic variability in pelagic
432 systems. In effect, the significance of ecological bridges and barriers will depend on the
prevalence of environmental events and the life history stage at which an individual exploits
434 a bridge or barrier.

436 For some of our case studies, the oceanographic drivers are unclear, but it is clear
437 that bridge and barrier dynamics can be influenced on a range of time scales – for example,
438 by decadal-scale cycles such as the El-Nino Southern Oscillation, the Pacific Decadal
439 Oscillation and the North Atlantic Oscillation (Higuchi et al., 1999). In ocean regions where
440 these climate drivers dominate, the biological responses may be influenced by the
441 appearance of bridges and barriers every few years (e.g. blue marlin). If there is a change
442 in the frequency of these climate modes (e.g. Table 1a), but no overall climate trend, then
443 the periodicity of the ecological bridge or barrier may also be affected. An increase in the
444 "breakdown" of the bridge conditions may lead to a decline in total population growth. Long-
445 term changes in bridge appearance (either declining or increasing frequency) have
446 occurred in the past, and are likely under climate change. Development of bridge
447 permanence, such as might be occurring in the Arctic now (e.g. bowhead whale), may lead
448 to loss of metapopulation structure if breeding between Atlantic and Pacific populations
449 (Table 1b), while a declining bridge frequency or barrier permanence may lead to great
450 population division and perhaps, over millennia, speciation (Table 1c). Ecological bridges
451 may be transient features in a changing climate, with the appearance of the bridge linked to
452 the rate of long-term change and the natural ocean variability.

452 **3.2 Population-level significance**

453 The significance of ecological bridges and barriers to pelagic species will likely be
454 dependent upon aspects of that species' life history characteristics and the ontogenetic
455 stage of individuals utilizing them. Important considerations include spatial range,
456 distribution, fundamental niche width, fidelity to breeding or foraging areas, and the relative
457 importance of proximate environmental influences versus learning and memory on at-sea
458 space use. Scaling from individual movements to population-level significance, ecological
459 bridges may be more readily exploited by neonate and juvenile stages of pelagic
460 organisms, as they disperse away from sites of natal origin.

461 In addition to the extent of a population's space use, the width of the fundamental
462 niche of a particular species may influence their propensity to use ecological bridges. A
463 recent theoretical model (Mariani et al., 2016) suggests that habitat suitability, migration
464 cost, and population structure can regulate habitat selection of highly migratory species.
465 Our case studies describe the broad ecological niche of bluefin tuna, which are able to
466 exploit a variety of prey types and tolerate a wide range of abiotic conditions (Arrizabalaga
467 et al., 2015; Fromentin et al., 2014a), and so can expand into novel regions with ease. More
468 specialised foragers, such as some surface-seizing and plunge-diving seabirds, require a
469 particular set of biophysical conditions and availability of certain prey types for effective
470 foraging, and so may be less likely to use ecological bridges in which conditions are not
471 energetically favourable (Ancona et al., 2012). For many species, particularly marine
472 ectotherms such as sea turtles, thermal sensitivity is a particularly important aspect that
473 might influence the response to barriers or bridges (Hawkes et al., 2007; McMahon and
474 Hays, 2006).

475 Moreover, fidelity to breeding and foraging habitats, and to migratory routes
476 between these habitats, is important to consider when questioning how movements through
477 ecological bridges might scale from individual- to population-level. A taxonomically diverse
478 range of marine vertebrates are known to demonstrate fidelity to particular foraging or
479 breeding habitats (e.g. tuna, Rooker et al. (2008); sharks, Queiroz et al. (2016); sea turtles,
480 Broderick et al. (2007); seabirds, Weimerskirch (2007)). This implies a considerable
481 influence on learning and memory in space use by a range of taxa (Regular et al., 2013).

484 Those that rely on learning and memory to navigate over proximate cues are less likely to
expand their range into new regions through an ecological bridge (Carroll et al., 2015). For
486 some fish species, it has been shown that the breakdown of information flow in a fish
community can cause habitat contraction and drive stocks to collapse in certain regions
488 (Petitgas et al., 2010). Moreover, theoretical analyses (Berdahl et al., 2016; De Luca et al.,
2014) suggest that for species moving in large groups (i.e. schooling), group formation can
490 be subject to threshold effects that alter migrations. For example, changes in individual
preference and/or of the total population density can produce rapid alterations in group
492 formation and collective behaviour to a point at which migration to other habitats may be
stopped (De Luca et al., 2014). As a result, any consideration of range expansion or
494 contraction must recognise the inherent interplay between an animal's responses to the
contemporaneous environment and the intrinsic motivations that underlie movements and
behaviours (Carroll et al., 2015).

496 In addition to species-specific constraints, ontogenetic stage may be important when
considering the significance of ecological bridges. While large pelagic fish are most readily
498 tagged, the movements of smaller juveniles and neonates may be of particular relevance to
ecological bridges. Individuals in dispersive life stages are more likely to expand the
500 population range into new habitats, as larval stages or neonates can often be advected in
prevailing current flow (e.g. sea turtle hatchlings, Hays et al. (2010)), and juveniles are more
502 likely to make exploratory movements at the edges of a population's current range (e.g. reef
sharks, Chin et al. (2013); breeding colony prospecting in immature seabirds, Dittmann et
504 al. (2005); Northern gannet on Farallon Islands, McKeon et al. (2015)). Conversely,
individuals of breeding age may be less likely to exploit opportunities resulting from
506 ecological bridges because many species show fidelity to particular breeding grounds, or
natal philopatry (e.g. turtles, Luschi et al. (1998); whales, Wedekin et al. (2010); sharks,
508 Feldheim et al. (2014); tuna, Block et al. (2005)).

The question of whether the significance of ecological bridges scales from changes
510 in individual movements over intra- to inter-annual timescales to population-level effects
remains unanswered. For some species, such as seabirds and marine mammals, that are
512 now able to move through the ice-free Northwest Passage, this novel connectivity between
habitats is almost certain to entail population-level effects, including genetic mixing, the
514 establishment of new breeding colonies for seabirds, and possible population expansion
into regions that marine mammals historically occupied but were extirpated. Thus,
516 connectivity can contribute to meta-population recovery of historically over-exploited
species, including various populations of marine mammals and bluefin tunas. This could
518 improve the resilience and sustainability of tuna fisheries, provided both tuna populations
and fisheries can adapt to novel spatio-temporal dynamics.

520

3.3 Future research

522 Advances in satellite telemetry and species distribution models have provided a
wealth of information linking the movements and behaviors of highly migratory species to
524 environmental conditions (e.g. Block et al. (2011); Hammerschlag et al. (2011); Hazen et al.
(2013); Hobday et al. (2011)). Integration of these findings reveals the importance of spatio-
526 temporal scales in understanding species-environment linkages (Hazen et al., 2013). Our
case studies describe changes in migratory corridors, which may be particularly important in
528 modifying the spatial dynamics of habitat use by populations of highly migratory species,
affecting circumpolar, trans-equatorial and trans-oceanic species distributions. However,
530 ecological bridges and barriers are likely to manifest over a range of spatio-temporal scales,

532 and further research into the mechanisms of biophysical coupling in the pelagic ecosystem
is necessary to truly understand the wider significance of these anomalous events. The
534 examples presented here are based on tracking and habitat models, yet limitations still exist
in our ability to track individuals throughout their life history stages and thus over the
536 environmental conditions experienced over a lifetime. Such information, coupled with
spatially explicit demographic models, may assist scientists and managers in developing
538 predictions and projections of species' responses to anticipated environmental change
(Dunning Jr et al., 1995).

540 While ecological bridges and barriers may be transient features in a changing
climate, they can foreshadow changes that are expected under climate change as
542 environmental tolerances are exceeded in some regions (Burrows et al., 2014). High-
resolution climate predictions (e.g. Popova et al. (2016)), may add further understanding as
544 to when, where, and how frequently bridges and barriers are likely to form over a variety of
spatial and temporal scales. Together, such models can be used to simulate changing
546 pelagic seascapes, providing management with scenarios to consider should an ecological
bridge or barrier originate, decline, or persist.

548

3.6 Conclusions

550 Understanding changes in the marine environment continues to be challenging.
Highly migratory species must navigate a fluid and shifting environment, adding complexity
552 to how behavioral adaptations occur in relation to their immediate environment. Here, we
have shown how ecological bridges and barriers can result in changes in highly mobile
554 species distributions, population dynamics, and connectivity with their proximate
environment. The availability of novel habitats through ecological bridges or disappearance
556 of traditional habitats through ecological barriers may impact on a range of pelagic species.
Important considerations include integration of life history characteristics and population-
558 level sensitivity to their environment, as well as a greater awareness and understanding of
the periodicity and frequency of bridges and barriers with time. As the effects of climate-
560 mediated ecosystem change are likely to be even more pronounced in the coming decades
(Burrows et al., 2014; Moore and Huntington, 2008), understanding how highly migratory
562 species navigate a changing environment will be more important than ever.

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572

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Table and Fig. Captions

902 **Table 1.** Schematic illustration of oceanic systems with bridge and barrier conditions over
904 time. Shaded grey area indicates time periods when the periodicity has changed in A, and
where bridges and barriers occur in B and C.

906 **Fig. 1.** (a) Examples of ecological bridges and barriers for five populations of pelagic
908 species along their migratory routes (solid white line). Dashed lines represent individual
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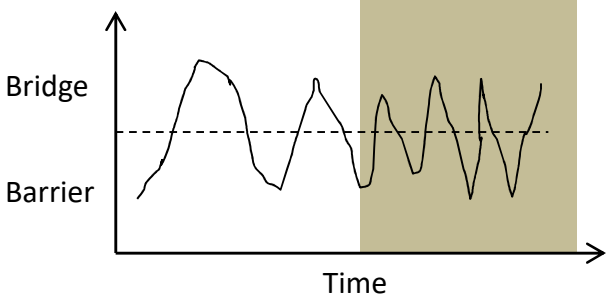
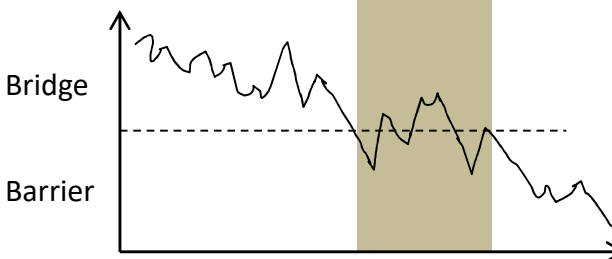
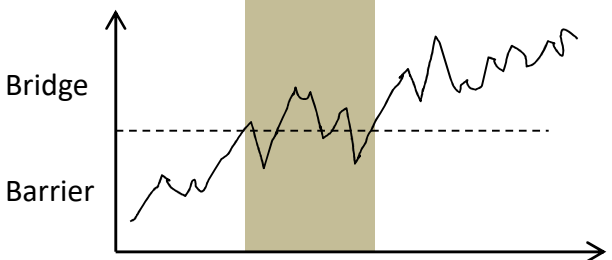
912 **Fig. 2.** Probabilities of Atlantic Bluefin tuna (ABFT) occurrence deduced from the NPPEN
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as the map of ABFT occurrence over 1960 to 1967 minus the median probabilities
916 calculated in each pixel from 1960 to 2009); and (c) same as (b) for the period 2000 to 2009
(from Fromentin et al., 2014a)

918 **Fig. 3:** Juvenile Atlantic Bluefin tuna potential feeding habitat (expressed as frequency of
920 occurrence) during migration period after wintering (from April to June) in 2009 (panel a),
2010 (panel b) and 2011 (panel c). The potential habitat of juvenile Atlantic bluefin tuna is
922 derived from the daily detection of chlorophyll-a fronts and a tolerance to sea surface
temperature (see Druon et al. 2016 for more details). Blanks indicate a frequency of
924 occurrence lower than 1%. The 200 m depth contour is shown. Panel d) shows the mean
926 occurrence of juvenile bluefin tuna feeding habitat in the central area of the bridge,
represented by a box (36-48°N, 35-57°W) in panels a, b and c, from April to June in each of
the years.

928 **Fig. 4.** Habitat preferences for juvenile SBT based on sea surface temperature (°C) and
930 chlorophyll a (mg/m³; log scale) for the area 25-45°S, 80-180°E during April-June of 1998-
2000 (a; n=46 fish) and 2004-2006 (b; n=24 fish); only fish ≥ 85 cm were included for
932 consistency between the two periods. Preferences were calculated by comparing
environmental data where SBT were located with environmental data for the whole area
934 during the time period of interest. Values >1 indicate preferred habitat (i.e. conditions at
which fish are found in greater proportion than they occurred in the ocean) (see Basson et
936 al., 2012). The maps show areas around Tasmania containing preferred SBT habitat
(values >1) in April-June of 1998-2000 (c) and 2004-2006 (d), based on the habitat
938 preference model for the corresponding time period.

940 **Fig. 5:** Trans-equatorial movements of blue marlin during the 2009 El Nino (right) and 2010
La Nina (left). Remotely sensed sea surface temperature is from October 2009 and 2010.
942 Tracks from 2009 and 2010 are shown, with the thick black sections showing period of track
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944 during every year of tagging except for during the 2010 La Nina (left). From Carlisle et al.
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946 **Table 1.** Schematic illustration of oceanic systems with bridge and barrier conditions over
 948 time. Shaded grey area indicates time periods when the periodicity has changed in A, and
 950 where bridges and barriers occur in B and C.

Mode	Example	Implications
<p>A. Increasing periodicity of bridge conditions</p> 	<p>Blue marlin – El Niño conditions break bridge more often</p>	<p>Context-dependent</p>
<p>B. Declining frequency of bridge conditions</p> 	<p>NW passage in historical times Salmon shark in north Pacific¹ Swordfish in Pacific²</p>	<p>Decreased connectivity, increase in metapopulations and possible speciation</p>
<p>C. Increasing frequency of bridge conditions</p> 	<p>Bowhead whale – NW passage³</p>	<p>Increased connectivity across species range – loss of metapopulation structure</p>

References: ¹Weng et al. (2005), ²Reeb et al. (2000), ³(Heide-Jørgensen et al., 2012)

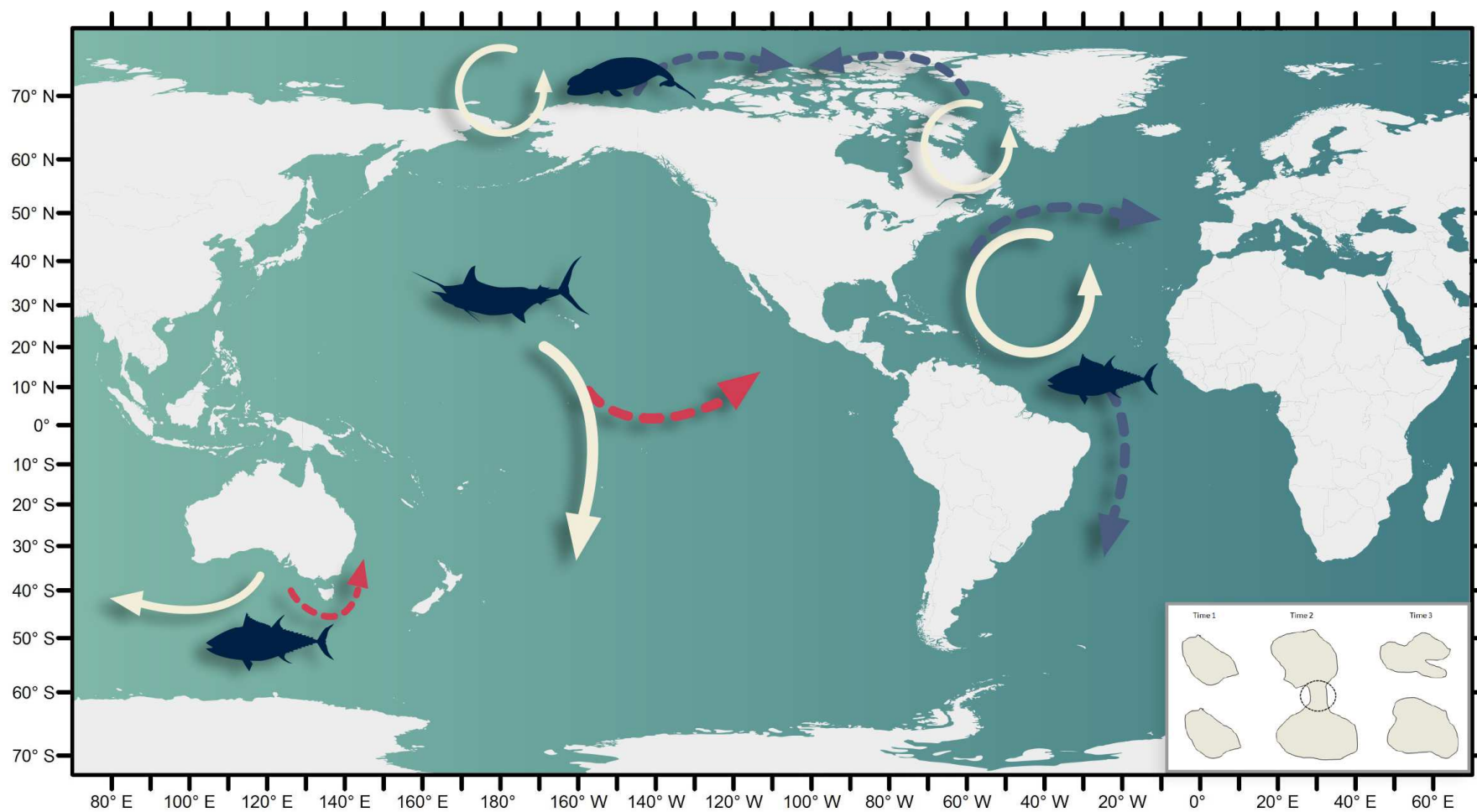
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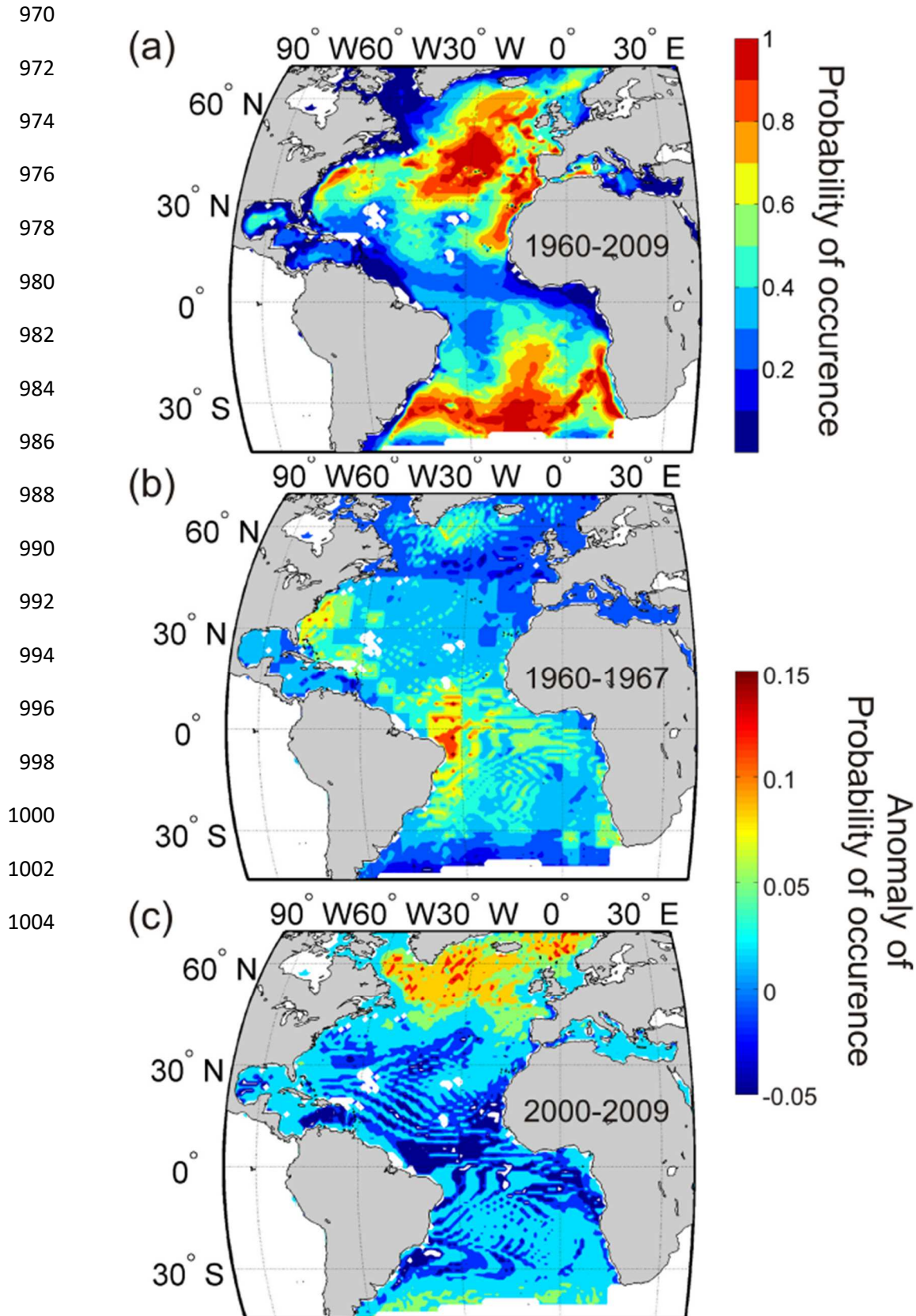
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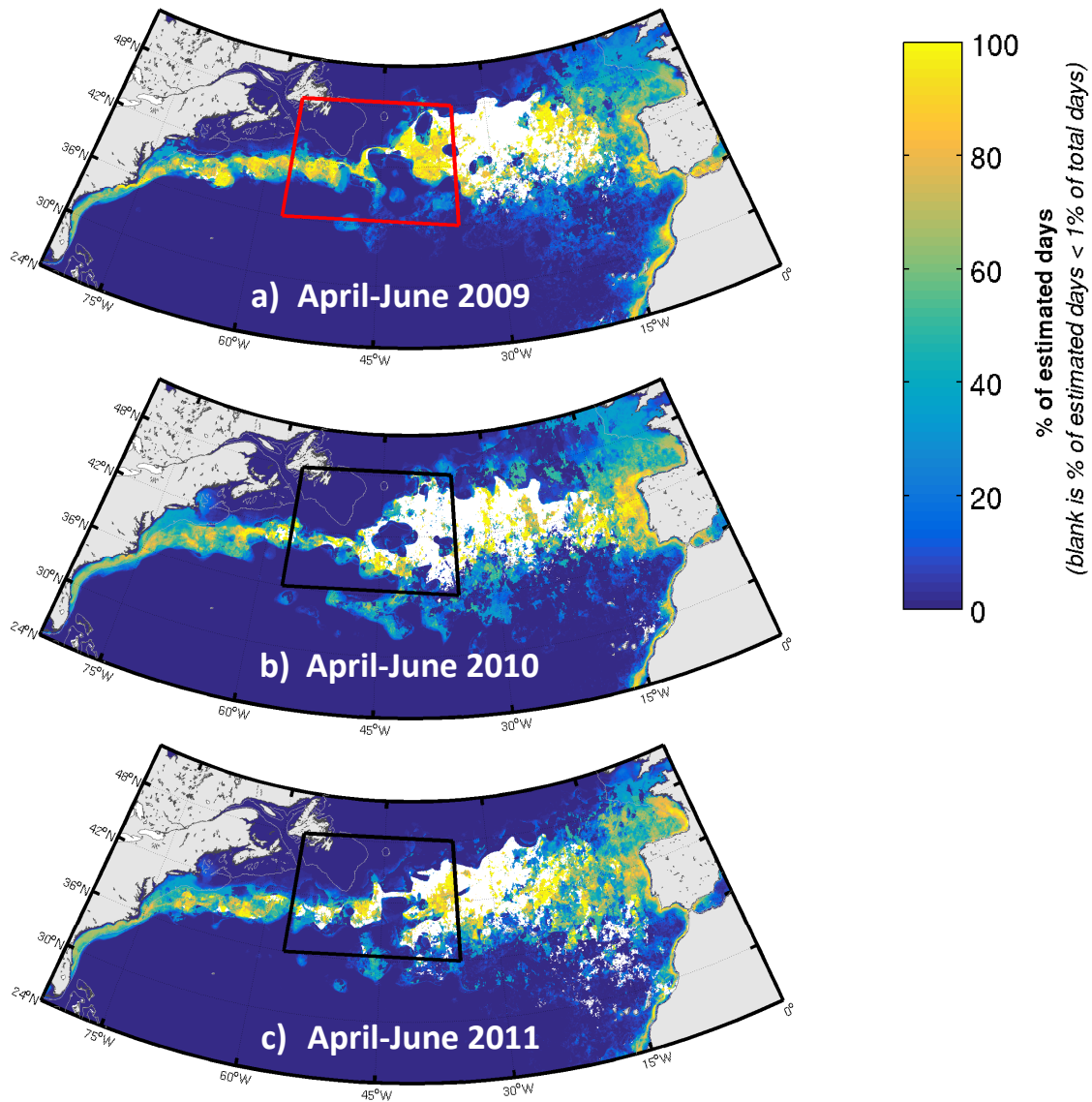
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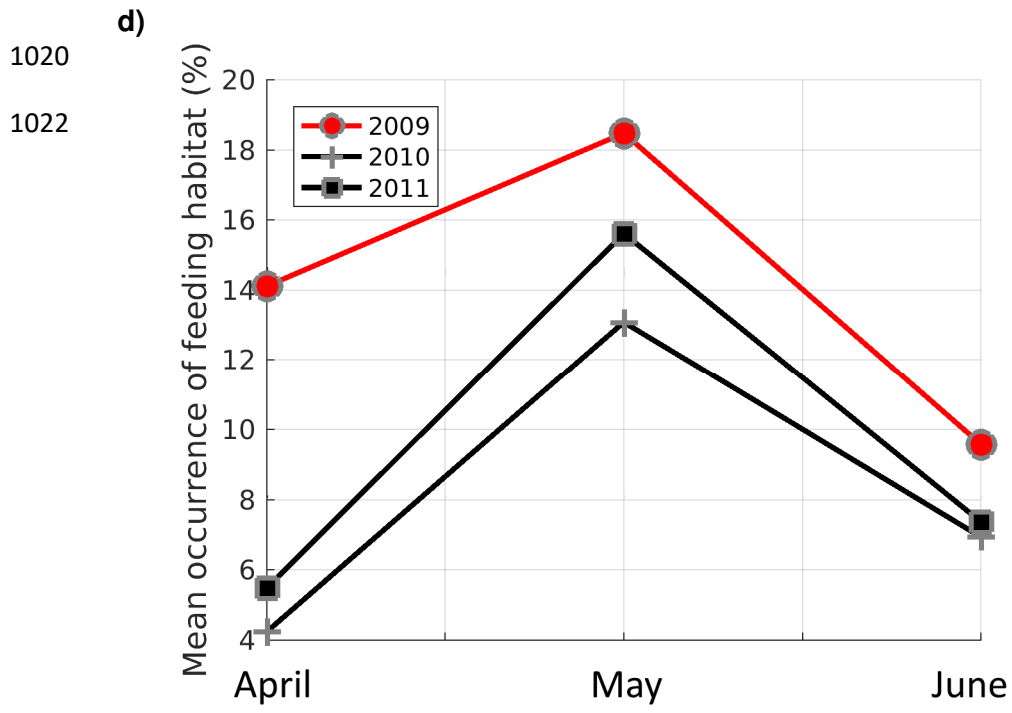


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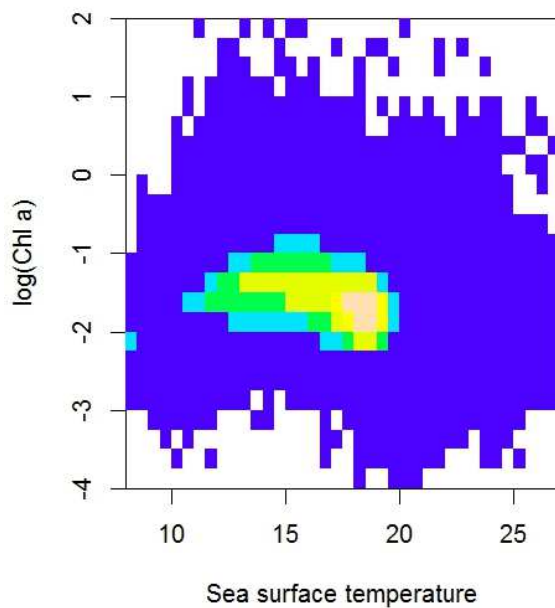
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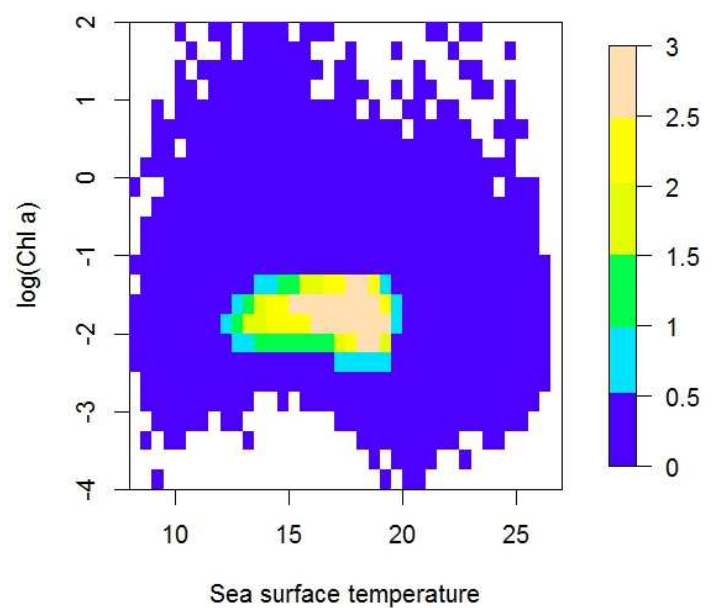
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(a)

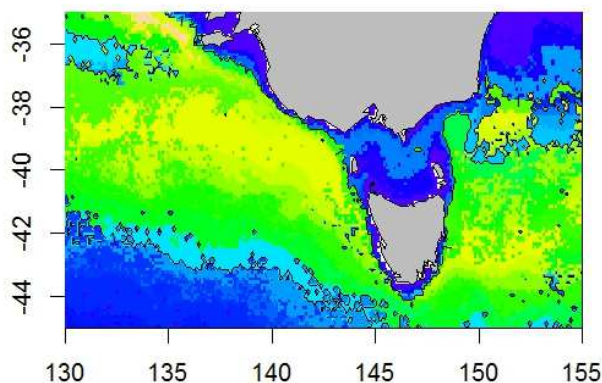


(b)

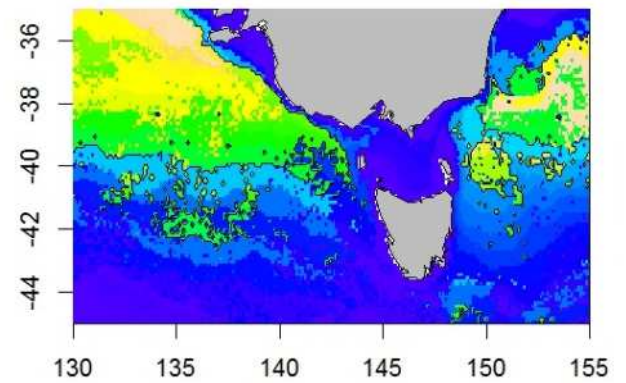


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(c)



(d)



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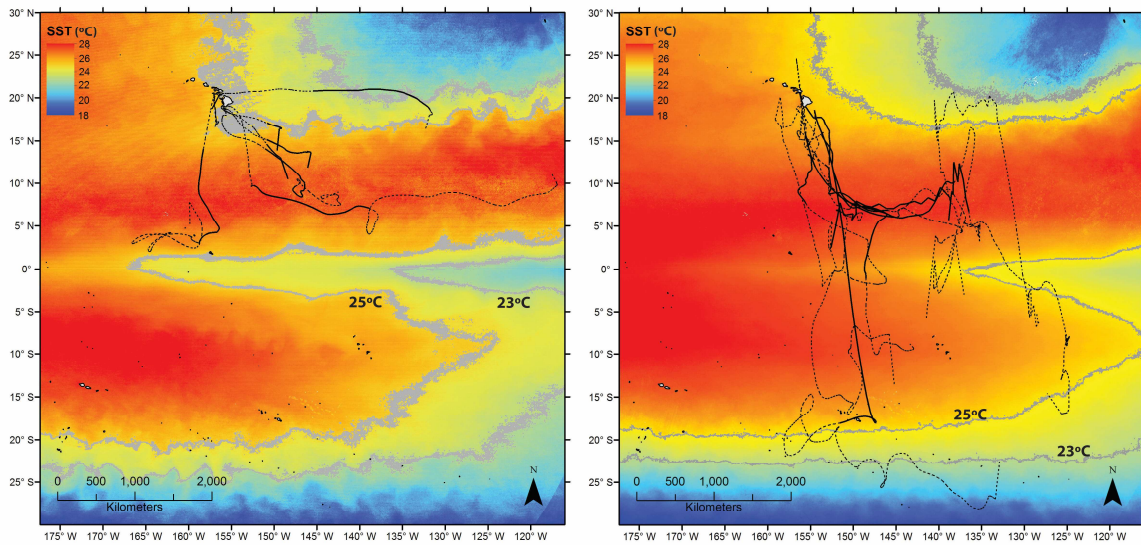
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