ABSTRACT: Interactions between seabirds and their prey are shaped largely by the dynamics of the marine ecosystems in which they are embedded. Physical oceanographic processes can drive the distributions of ectothermic and planktonic prey and hence their availability to marine birds. Owing to the complex nature of these processes and interactions, the signal-to-noise ratios of avian indicators of prey conditions are variable, often low and further degraded (buffered) by seabird behaviour and life-history features. Cairns (1987, Biol Oceanogr 5:261–271) detailed seabird responses operating over a range of temporal scales to variation in food supplies, and suggested that interval relationships might be developed between avian responses and environmental variation. While this may be possible in some instances, it appears unrealistic in most instances to expect interval relationships between seabird responses and prey conditions that are often nonlinearly related. The present paper focuses on binary data (e.g. breeding success versus failure) derived from seabirds that can provide robust information about major shifts in prey and oceanographic conditions and that are particularly informative when accumulated over decadal and large ocean scales. Inter-annual and decadal variations in specific and nominally categorized (warm- versus cold-water) prey landings of northern gannets *Sula bassana* at a large oceanic colony in the NW Atlantic reflect shifts in pelagic food webs induced by changes in regional sea surface temperature. Binary patterns emphasize decadal shifts in food webs and yield predictive indication of systemic change.

KEY WORDS: Binary patterns · Bio-indicators · Decadal scales · Seabirds · Ecosystem shift

INTRODUCTION

Considerable research has been directed at the exploitation of information from marine birds and mammals for purposes of assessing the changing states of prey bases and for detecting oceanographic and fisheries effects on top predators (e.g. Boyd et al. 2006, Piatt et al. 2007a, this issue). The physical dynamics of oceans exert considerable influence on interactions between seabird predators and their prey. Driven largely by ocean conditions, the distributions of ectothermic and planktonic prey determine their availability to avian predators (Scott et al. 2006). Owing to the stochastic nature of these processes and interactions, the signal-to-noise ratios of seabird indications of prey conditions are variable, frequently low and are degraded (buffered) by avian behaviour and life-history adaptations (e.g. flexible foraging tactics, deferred reproduction; Burger & Piatt 1990, Montevecchi & Berruti 1991, Daunt et al. 2006). Consequently, it is not simple to effectively derive insight from seabird data about changes in prey and ocean conditions. Yet biological signals complement and can even be more revealing than physical signals in detecting pervasive ocean perturbations, such as regime or ecosystem shifts (McFarlane et al. 2000, Hare & Mantua 2000).

In a seminal paper on bio-indicators, Cairns (1987) proposed patterns of seabird responses operating over a range of temporal (and hence spatial) scales to changes in their prey base. The responses considered
ranged from population through to behavioural levels and included adult survival, breeding success, chick growth, parental attendance and activity (foraging) budgets. Each response, with the exception of foraging budgets, was hypothesized to exhibit relatively sharp nonlinear changes to different levels of prey availability, which was considered on an ordinal scale between low and high. Cairns (1987, p 267) contended that ‘[a]t present seabird data can yield information on marine food supply on ordinal scales, but assigning food availability to interval or ratio scales must await rigorous testing of the relationships...’. Over the 20 yr since Cairns’ paper appeared, there have been numerous attempts to assess seabirds as indicators of prey conditions. The most recent of these sought to directly assess the predictions of Cairns’ (1987) models, and found support for some of the predictions and not for others (Piatt et al. 2007b, this issue).

However, in most instances it is unrealistic to derive interval scaling between seabird responses and prey conditions that are often nonlinearly related. Yet this circumstance in no way precludes the derivation of information about marine ecosystems from seabird research. Associations of nominal patterns of seabird biology and prey and oceanographic conditions have proven highly informative in many instances (e.g. Schreiber & Schreiber 1984). Binary relationships between avian predators, prey and oceanography gain considerable power and robustness as they are cumulated over time, space and species (Hatch 1996). For example, in an ocean-basin study of black-legged kittiwakes Rissa tridactyla, Hatch et al. (1993) demonstrated striking inter-annual patterns in the concordances and dis-concordances of reproductive success versus failure among colonies in the Bering Sea and Gulf of Alaska from 1978 through 1989 (Fig. 1). In 1978, colonies in the Bering Sea failed, while those in the Gulf of Alaska were productive. In 1984, this pattern reversed, whereas in 1988 all colonies throughout the study area were successful and in 1989 virtually all colonies failed. The information derived from these signals captured systemic patterns that occurred over scales that were larger than those of fishery activities and oceanographic research in the region.

The present paper builds on the potential of most studies of seabird ecology to detect binary changes in reproductive and behavioural parameters in response to environmental change. In this respect, I compare inter-annual and decadal associations in the species compositions of prey landings by northern gannets Sula bassana at a large oceanic colony with sea surface temperature (SST) patterns and anomalies in the NW Atlantic. Prey landings are assessed on a single-species basis and in nominal multi-species ecological classifications of prey from temperate/subtropical and from low Arctic Ocean regions, hereafter referred to as warm- and cold-water prey. Emphasis is directed at the cumulative information that can be derived for

![Fig. 1. Rissa tridactyla. Inter-annual ocean-basin patterns of breeding success (●) and failure (○) of black-legged kittiwakes at colonies in the North Pacific (after Hatch et al. 1993)](image-url)
these types of responses when combined over decades and longer and over large oceanographic scales to indicate shifts in food webs and ecosystem conditions.

**MATERIALS AND METHODS**

**Study site.** Research was conducted in Funk Island Ecological Reserve (49°45′N, 53°11′W), a small (800 × 400 m) flat granite rock in the northwest Atlantic (Montevecchi & Tuck 1987), about 50 km off the northeast Newfoundland coast (Fig. 2), Canada. About 10,000 pairs of northern gannets *Sula bassana* nest in the colony (Chardine 2000), which is the third largest and most oceanic of the 6 gannet colonies in North America.

**Prey load sampling.** Food samples were obtained by approaching roosting gannets that often regurgitated as they moved away from researchers (Montevecchi & Myers 1995). Some samples were also obtained when gannets had GPS and other data loggers attached and removed and from discarded regurgitations and scraps in the colony. While there are likely differences between samples collected in roosts and in the colony, samples from all sources were comparable, so most regurgitations were collected at roosts well outside the colony to minimize disturbance to nesting areas. Regurgitated prey were identified to species, sex and condition (gravid, spent/immature) whenever possible. Prey landings are presented as percentages of the total estimated mass of prey landed during each year. A total of 8239 prey samples were obtained during late July and early to mid-August from 1977 to 2006, with the exception of 1981, when it was not possible to land on the island (Table 1).

**Sea surface temperatures.** SSTs (0 to 20 m), corresponding with the maximum dive depths of northern gannets (Garthe et al. 2000), were obtained from Hydrographic Stn 27 (48°32.8′N, 52°35.2′W) located 250 km S and downstream of Funk Island in the Labrador Current. The station has been checked regularly since its establishment in 1946. Measurements from Stn 27 provide robust ocean climate signals for the entire Newfoundland–Labrador Shelf (Petrie et al. 1988, Myers et al. 1990). Annual averages of SST during June, July and August were used.

Table 1. *Sula bassana*. Dates of collection and number of prey samples from gannets in the colony on Funk Island, 1977 to 2005

<table>
<thead>
<tr>
<th>Year</th>
<th>Date</th>
<th>With 1 species</th>
<th>With &gt;1 species</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>1977</td>
<td>12 Jul</td>
<td>105</td>
<td>0</td>
<td>105</td>
</tr>
<tr>
<td>1978</td>
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<td>496</td>
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<td>508</td>
</tr>
<tr>
<td>1979</td>
<td>31 Jul–6 Aug</td>
<td>162</td>
<td>4</td>
<td>166</td>
</tr>
<tr>
<td>1980</td>
<td>1–9 Aug</td>
<td>217</td>
<td>7</td>
<td>224</td>
</tr>
<tr>
<td>1982</td>
<td>9–13 Aug</td>
<td>191</td>
<td>25</td>
<td>216</td>
</tr>
<tr>
<td>1983</td>
<td>5–13 Aug</td>
<td>488</td>
<td>2</td>
<td>490</td>
</tr>
<tr>
<td>1984</td>
<td>9–19 Aug</td>
<td>225</td>
<td>8</td>
<td>233</td>
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<td>1985</td>
<td>4–10 Sep</td>
<td>184</td>
<td>16</td>
<td>200</td>
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<td>1986</td>
<td>7–15 Aug</td>
<td>493</td>
<td>22</td>
<td>515</td>
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<td>561</td>
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<td>18 Jul–11 Aug</td>
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<tr>
<td>1991</td>
<td>13 Jul–16 Aug</td>
<td>411</td>
<td>21</td>
<td>432</td>
</tr>
<tr>
<td>1992</td>
<td>5–10 Aug</td>
<td>293</td>
<td>8</td>
<td>301</td>
</tr>
<tr>
<td>1993</td>
<td>5–12 Aug</td>
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<tr>
<td>1994</td>
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<td>1995</td>
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<tr>
<td>1996</td>
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<td>1997</td>
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<td>244</td>
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<td>246</td>
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<td>1998</td>
<td>3–17 Aug</td>
<td>446</td>
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<td>460</td>
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<tr>
<td>2003</td>
<td>26 Jul–6 Aug</td>
<td>391</td>
<td>2</td>
<td>393</td>
</tr>
<tr>
<td>2004</td>
<td>Jul 26–Aug 1</td>
<td>88</td>
<td>1</td>
<td>89</td>
</tr>
<tr>
<td>2005</td>
<td>3–11 Aug</td>
<td>200</td>
<td>0</td>
<td>200</td>
</tr>
<tr>
<td>2006</td>
<td>6–14 Aug</td>
<td>216</td>
<td>7</td>
<td>223</td>
</tr>
<tr>
<td>Totals 1977–2006</td>
<td>7932</td>
<td>307</td>
<td>8239</td>
<td></td>
</tr>
</tbody>
</table>
Data analyses. One-way ANOVAs and Tukey post hoc tests were used to compare decadal differences in the prey landings of gannets; chi-squared and binomial tests were used to assess decadal binary patterns in the gannets’ landings of warm- and cold-water prey (Siegel 1957). Pearson correlations were run between individual prey species and between amalgamations of warm- and cold-water prey landings and SST. Level of significance is taken as p < 0.05.

RESULTS

Inter-annual variation in prey landings

As evident in their prey landings at Funk Island since 1977, gannets are generalist predators that exploit a diverse array of pelagic fish and squid with considerable inter-annual variation (chi-squared = 10 801.6, df = 130, p < 0.0001; Fig. 3). From 1977 to 1980, 77 ± 11% of the prey landed was Atlantic mackerel *Scomber scombrus*. In 1982, mackerel made up only 13% of the estimated prey landed and was replaced by Atlantic herring *Clupea harengus*, which represented 60% of the prey landed and which is second to mackerel in energy density among the gannets’ prey spectrum (Montevecchi et al. 1984). In 1983, the relationship between the landings of these pelagic fishes was reversed, with mackerel comprising 83% of the landings and herring 12%; this relationship of greater landings of mackerel compared to herring continued until 1989. From 1989 onward, mackerel was landed very irregularly by the gannets, with no landings from 1996 through 2001 and with contrasting large landings in 2005 and 2006. Her- ring were landed in large proportions in 1993 and 1994 (35 and 37% of total mass, respectively; Fig. 3).

From 1977 to 1982, short-finned squid *Illex illecebrosus* made up from 7 to 21% of the prey landed, averaging 11 ± 6% per yr. In contrast, squid were only landed in 7 of the years between 1983 and 2006 (1989 to 1992, 1996, 1997, 2004), comprising 5% or usually less of the prey.

From 1977 to 1983, Atlantic saury *Scomberesox saurus* were either not landed (1977) or were a very minor component of the landings, ranging from 1 to 10%. Then, in the mid- to late 1980s, saury was a major component of the gannets’ prey every year except 1986 (percent of total mass of prey landed = 54% [1984], 39% [1985], 4% [1986], 19% [1987], 44% [1988], 63% [1989]). From 1990 to 2004, no saury were landed in most years, in 5 of the years <1 to 3% were landed, and saury comprised 12 and 13% of the prey landed in 1994 and 1998, respectively. In 2005 and 2006, 33 and 80% of the prey landed by the gannets were Atlantic saury (Fig. 3). The mean percentages of saury landed during the 1970s/1980s (20.85 ± 22.95), 1990s (2.65 ± 4.78) and 2000s (16.66 ± 30.44), though differing by an order of magnitude, are not significantly different owing to the high inter-annual variability associated with the means ($F = 2.15$, df = 2, 26, p = 0.136).

Landings of Atlantic salmon *Salmo salar* during the 1970s/1980s, 1990s and 2000s differed significantly ($F = 3.87$, df = 2, 26, p = 0.034). From 1977 to 1989, Atlantic salmon was landed in minute amounts, ranging from 0 to 1.4% of the estimated total landings, averaging 0.31% of the prey landed per yr. Circumstances changed in 1990, and from 1990 to 1996 salmon comprised from 3 to 7% of gannets’ annual prey landings. During the 1990s, Atlantic salmon averaged 3.4% of the total prey landed per yr. The largest harvests of salmon occurred in 2001 and 2002 (27 and 32% of the total mass of prey landed, respectively). Seven percent of the landings in 2003 were salmon, and none were landed thereafter (Fig. 4). Landings from 2000 to 2006 averaged 9.3% of the annual prey landings.

From 1977 to 1989, capelin *Mallotus villosus*, a small forage fish, comprised <5% of the mass of gannets’ prey landings in all years except in 1978 (9%) and 1982 (12%), when small amounts were also landed (Fig. 3). From 1989 until 2004, capelin was a common prey of gannets, ranging from 26 to 100% of prey landed each year. Significantly less capelin was landed by gannets...
in the 1970s/1980s than in the 1990s and 2000s ($F = 23.68$, $df = 2, 26$, $p < 0.001$ and Tukey post hoc test).

Small amounts of cod *Gadus morhoa* (discards) and sandlance *Ammodytes* sp. were also landed in 1 or 2 of the years.

### Decadal variation in prey landings

When these highly variable prey landings of the gannets are collapsed into the binary categories of warm-water migratory prey (mackerel, squid, saury) from temperate and cool subtropical ocean regions and cold-water low arctic prey (herring, capelin, salmon, sandlance, cod), striking decadal patterns emerge (Fig. 5). Migratory warm-water species comprised most of the gannets' prey landings in the late 1970s and throughout the 1980s (binomial test of equal proportions of warm- and cold-water prey, $p = 0.003$). In a marked reversal of this pattern, from 1990 through 2004 cold-water prey comprised almost all of the gannets' prey landings (binomial test of equal proportions of warm- and cold-water prey, $p < 0.001$). A comparison of the landings of warm-water migratory prey landed before and after the 1991 cold-water perturbation (Fig. 6) proves to be highly significant ($F = 43.57$, $df = 1, 27$, $p < 0.001$). In 2005 and 2006, for the first time since 1989, migratory warm-water pelagic fishes comprised 81 and 93%, respectively, of the estimated prey landed by gannets.

### Annual and decadal associations between prey landings and SST

Following the lowest SST on record in 1991, values returned to positive thermal anomalies by the mid- to late 1990s (Fig. 6). Still, warm-water migratory prey did not regain their dominance in the gannets’ prey landings until 2005, lagging about a decade after the reoccurrence of positive SST signals (Fig. 4). Sixteen years passed (from 1989 to 2005) before warm-water prey again dominated landings. 2005 was a warm-water year in the region (but not the warmest in recent years). There were no significant associations between the annual landings of any prey or combined landings of warm- or cold-water prey and average annual SST from 1977 to 2006. The prevalence of migratory warm-water pelagic species in the gannets’ prey landings continued in 2006.
DISCUSSION

A highly flexible foraging strategy directed at a wide breadth of pelagic prey is a striking feature of the northern gannets' *Sula bassana* feeding ecology (e.g. Garthe et al. 2007). The generalist pattern of their changing dietary diversity is evident in their prey landings on Funk Island from 1977 through 2006. Inter-annual variation in the gannets' landings of mackerel and squid exhibited significant correlations with commercial landings and independent research indices of these species over multiple spatial and temporal scales beyond the gannets' foraging range around the colony (Montevecchi & Myers 1995). These associations are driven by prey availability, largely by negative events when birds and humans catch little or none of a particular pelagic species.

### Physical forcing and lagging biological responses

The 1991 cold-water perturbation in the NW Atlantic—a centennially significant event (Drinkwater 1996)—acted as a physical forcer that influenced and inhibited warm-water pelagic species such as mackerel, saury and squid from migrating into the region (Montevecchi & Myers 1995). This cold-water incursion induced an extensive regime-type shift in the pelagic food web (Montevecchi & Myers 1996). By the mid-1990s, SSTs had returned to pre-perturbation levels, yet a return to prior warm-water prey landings by both birds and commercial fishers lagged by about another 10 yr (see also Davoren & Montevecchi 2003). It is this type of on/off response that is responsible for the lack of inter-annual associations (correlations) between the prey landings of birds and average annual SSTs. These decadal patterns and their coincidence with oceanographic perturbation are clarified when the variation in inter-annual prey landings are lumped into the binary categories of warm- and cold-water prey.

### Binary responses and systemic shifts

Binary classification, amalgamation and analysis can provide robust signals from apex marine predators (see Hatch 1996 for analyses of binary patterns). In 2005, the prominence of migratory warm-water mackerel and saury in the gannets’ landings returned for the first time in 16 yr. This avian signal of warm-water prey also coincided with the largest mackerel fishery landings in the Newfoundland region since 1989 (NAFO Division 3K and 3L reports; M. Koen-Alonso pers. comm.). The changes in prey landings by the gannets signalled a shift in the pelagic food web driven by oceanographic changes, leading to greater representation of migratory warm-water pelagic species in regional seasonal assemblages and food webs. These signals from seabirds can also be used to anticipate predator–prey and wider food web and oceanographic patterns. As predicted (Montevecchi 2005), ocean temperatures were warm and the mackerel fishery flourished in 2006. The prey landings of gannets in 2006 were again dominated by migratory warm-water prey from temperate and subtropical ocean regions, and ocean temperatures in the NW Atlantic were among the warmest on record (E. Colbourne pers. comm.). In the NW Atlantic in 2006, auks laid their eggs about 2 wk earlier than in previous years, spawning capelin and humpback whales *Megaptera novaeanglia* moved inshore early, lumpfish *Cyclopterus lumpus* were egg-laden in May, well before their usual time during summer and large squid were abundant in inshore waters (W.A.M., D. Fifield, A. Hedd, J. Lavers, T. and K. Power pers. obs.). So at least during 2005 and 2006, a shift back to warm-water ecosystem conditions was evident. Continuation of this warm-water condition occurred in 2007 and is expected to continue further.

### Seabird indicators

Seabird studies of biophysical environmental conditions have proven highly informative in other contexts. For instance, broad-scale biological shifts are driven at times by radical state changes (abundance versus scarcity) exhibited by focal forage species that fuel large vertebrate food webs (Chavez et al. 2003). Seabird studies are particularly useful in documenting condition changes in these forage species (e.g. Österblom et al. 2001, Davoren & Montevecchi 2003, Miller & Sydeman 2004, Wanless et al. 2004, 2005) that often reflect food resource states. Importantly, seabirds also target prey that are not commercially exploited (and hence excluded from most fishery research programs) or are not accessible to standard survey methods (Barrett et al. 1990, Montevecchi 1993). Recent studies use data-logging devices to interrogate free-ranging seabirds and mammals about environmental information (Wilson et al. 2002, Daunt et al. 2003, 2006, Garthe et al. 2007). When carried out in conjunction with vessel surveys of prey densities and distributions, these studies hold the further potential of detailing the behaviour and decision-making of individual foragers (Garthe et al. 2000, Ollason et al. 2006, Staniland et al. 2006) and directly assessing functional responses, the mechanisms of higher level population patterns.
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