The decline of Norwegian kittiwake populations: modelling the role of ocean warming

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ABSTRACT: The black-legged kittiwake Rissa tridactyla is a pelagic seabird whose population has recently declined in most parts of the North Atlantic and which is red-listed in most bordering countries. To investigate a possible cause for this decline, we analysed the population dynamics of 5 kittiwake colonies along the Norwegian coast, ranging from 62° to 71° N, over the last 20 to 35 yr. By quantifying the importance of sea surface temperatures (SST) in relevant areas of the North Atlantic, we tested the importance of climatic conditions throughout the populations’ annual cycles. We found no synchrony among colonies; however, SST affected population dynamics, explaining between 6% and 37% (average 18%) of the variation in annual population growth rate. While dynamics of the southerly colonies were mainly affected by winter conditions in the Grand Banks area, dynamics of the northerly colonies were dominated by autumn conditions off Svalbard. Negative slopes indicated stronger population decline under warmer ocean conditions. Population dynamics were affected both via adult survival and offspring recruitment, as evidenced by the presence of unlagged effects as well as effects lagged by the age at recruitment. Finally, we performed population viability analyses taking into account the projected warming trends for the future. The median time to extinction of the Norwegian colonies was 52 to 181 yr without considering covariates; 45 to 94 yr when considering the effects of SST but ignoring future warming; and 10 to 48 yr when ocean warming, based on a ‘business as usual’ scenario, was taken into account.

KEY WORDS: Global warming · Non-breeding distribution · Population dynamics · Population viability analysis · Rissa tridactyla · Sea surface temperature

1. INTRODUCTION

Predicting the impact of human activities, including global climate change, on the biosphere has become one of the most important efforts in ecology. Ecosystems worldwide are changing rapidly as a consequence of anthropogenic impacts such as global warming (IPCC 2007), yet our understanding of the consequences of these changes on populations is limited. To be able to predict population trajectories, it is crucial to understand the mechanisms underlying variation of, and co-variation among, populations. Population fluctuations are determined by parameters such as intrinsic population growth rate and carrying capacity, as well as by stochastic fluctuations in the environment (Lande et al. 2003). Furthermore, temporal variation in climate and other environmental variables may synchronise population
fluctuations over large distances (Moran 1953, Bjørnstad et al. 1999, Lande et al. 1999, Post & Forchhammer 2002). Population synchrony, defined as the inter-annual correlation of population growth rates across colonies, can therefore indicate the presence of environmental factors affecting population dynamics on large spatial scales. Moreover, a high degree of inter-annual synchrony in population fluctuations increases the risk of local and global extinction (Esler 2000, Engen et al. 2002). Herein lies the main importance of population synchrony for population management.

In seabirds, knowledge about their distribution outside the breeding season has for a long time been a limiting factor in analysing and understanding the importance of environmental conditions for population processes (e.g. Smith & Gaston 2012). Seabirds are normally philopatric and return to the same breeding colony each year, but may disperse over vast ocean ranges during the rest of the year (e.g. Egevang et al. 2010, Frederiksen et al. 2012). Quantitative analyses of environmental conditions during non-breeding have only recently become feasible through advances in tracking technologies, such as miniaturized year-round light-based tracking devices (GLS loggers or geolocators; Phillips et al. 2004, González-Solís et al. 2007, Egevang et al. 2010, Seavy et al. 2012).

We here use the novel knowledge of non-breeding distribution (Frederiksen et al. 2012) to search for environmental covariates explaining population dynamics of the black-legged kittiwake *Rissa tridactyla* (hereafter called kittiwake) in Norway. Kittiwake numbers have declined over most of the North Atlantic over the last 2 decades, particularly in the North Sea and adjacent areas (Frederiksen 2010). According to the IUCN Red List, the species is of least concern globally; in national Red Lists, however, it is currently listed as near threatened in Denmark, France and Svalbard, as vulnerable (or ‘amber’) in the Faroes, Greenland, Great Britain and Ireland and as endangered in Norway and Sweden (Wind & Pihl 2004, Fosaa et al. 2005, Boertman 2007, Lønnes et al. 2007, Eaton et al. 2009, Gårdfors et al., 2010, Kålås et al. 2010, UICN France et al. 2011).

At a study colony in the North Sea, the decline was caused by low reproductive success as well as low adult survival (Frederiksen et al. 2004), both apparently linked to increasing sea temperatures affecting their main prey (Frederiksen et al. 2006). To understand whether the ocean-wide decline of kittiwakes is governed by a common factor, it is important to establish whether these findings can be generalised to kittiwake populations in other areas.

We here study the dynamics of 5 kittiwake populations along the coast of Norway in order to address 4 questions: (1) Is there any population synchrony between the colonies? (2) How are the population dynamics related to the local climatic conditions (sea surface temperature [SST]) in the areas where the birds stay during different parts of their annual cycle? (3) Through which demographic trait (adult survival or offspring recruitment) is climate affecting the population growth rate? (4) How will the predicted future warming trend affect the viability of the populations?

### 2. MATERIAL AND METHODS

#### 2.1. Population monitoring

Kittiwakes breed in many places along the Norwegian coast. We analysed population counts from the 5 kittiwake *Rissa tridactyla* colonies that are part of the long-term Norwegian Monitoring Programme for Seabirds, covering the geographic range from the southern Norwegian Sea to the Barents Sea (Fig. 1). From southwest (boreal climate) to northeast (Arctic climate), the colonies included were Runde (62° 24’ N, 5° 38’ E), Sklinna (65° 12’ N, 10° 59’ E), Vedøy (67° 29’ N, 12° 1’ E), Hjelmsøya (71° 4’ N, 24° 43’ E) and Hornøya (70° 23’ N, 31° 9’ E).

The populations were monitored according to standardised methods (e.g. Walsh et al. 1995) using apparently occupied nests or nest sites (AON) as the counting unit. At Sklinna, the whole colony was counted, while averaged counts in randomly selected study plots were used in the other colonies. Annual estimates of AONs were based on a total count made in 2010/2011 and the annual rates of change documented in the monitoring plots. The AONs in the study plots represented 1.1% (10 study plots), 100% (whole colony count), 3.7% (6 study plots), 5.2% (5 study plots) and 12.3% (9 study plots) of the total population in Runde, Sklinna, Vedøy, Hjelmsøya and Hornøya, respectively. All counts were made late in the incubation period or early in the chick period.

Colony sizes varied by 3 orders of magnitude among colonies, ranging from 170 pairs in Sklinna to 158 000 in Runde in 1980. All 5 populations declined during the monitoring period that spanned 20 to 35 yr (Fig. 2). The breeding population at Sklinna went extinct in 2011; in our models, we disregarded all counts at this colony after 2001, when population size for the first time dropped below 20 breeding pairs.
2.2. Population models

Population dynamics of the 5 colonies were density-independent, as evidenced by the absence of any negative correlation between annual growth rates $r_t$ and population sizes $N_t$ (all correlation coefficients $R > -0.25$, all $p > 0.3$; see the Supplement at www.int-res.com/articles/suppl/c061p091_supp.pdf for density-dependent models). We therefore used Brownian population models of the following form:

$$\ln N_{t+1} = \ln N_t + \bar{r} - \frac{1}{2} \sigma_d^2 N_t + \sum \beta_i X_{ij} + \epsilon_t$$  \hspace{1cm} (1)

with $\beta_i$ as the slope of the $i$th environmental covariate $X_{ij}$; $\epsilon$ is the environmental noise, i.e. an independent variable with zero mean and variance $\sigma_e^2$ (environmental variance); $N_t$ is the population size in year $t$; $\bar{r}$ is the long-term intrinsic population growth rate; $\sigma_d^2$ is the demographic variance; $X_{ij}$ is the environmental covariate $i$ in year $t$. The parameters $\beta_i$, $\bar{r}$ and $\sigma_e^2$ were estimated from the population time series using maximum likelihood such that the log-likelihood

$$\ln L = -\frac{1}{2} \sum_{k=2}^n \left[ \ln N_k - E(\ln N_k) \right]^2 \sigma_e^{-2} + \ln(2\pi \sigma_e^2)$$  \hspace{1cm} (2)

was maximised over the $n$ elements of the time series (Sæther et al. 2009), where $E(\ln N_k)$ is the predicted log-population size based on the observed population size $N_{k-1}$ and Eq. (1), and $\sigma_e^2 = \sigma_d^2 + \sigma_i^2/N$. In the absence of estimates of life-time reproductive success of the 5 colonies, demographic variance was assumed to be 0.1 in all colonies, which is a realistic value for long-lived birds (Lande et al. 2003).

Population models were either fitted to one colony at a time or to all populations simultaneously. In the former case, the optimal set of parameters for each colony could be identified. The latter approach enabled us to test for the presence of synchrony and whether population parameters differed among colonies. It was carried out by modifying Eq. (1) in such a way that any of the parameters $\bar{r}$, $\sigma_e^2$ or $\beta_i$ could be replaced by a vector of length 5, containing the growth rates, environmental variances or slopes for each of the 5 colonies; log-likelihood was maximised over the $\sum_{t=1}^5 n_t$ elements of all 5 time series using Eq. (2). These modifications could be combined such that, for example, models with a common $\bar{r}$ and $\sigma_e^2$, with a common $\bar{r}$ and 5 separate $\sigma_e^2$, with a common $\sigma_e^2$ and 5 separate $\bar{r}$, and with 5 separate $\bar{r}$ and $\sigma_e^2$ could be compared with each other (likewise for each of the environmental variables $\beta_i$). If a model with a common parameter was preferred over a model with separate parameters, this indicated that this specific parameter did not differ significantly among colonies. Specifically, a common estimate for the slope $\beta$ would indicate that the corresponding environmental covariate is common to all 5 colonies and synchronises their dynamics.

Models with different parameterisations or covariates were compared using Akaike’s information criterion corrected for small sample sizes ($AIC_C$), preferring models with the lowest $\Delta AIC_C$ (or the highest $AIC_C$ weight or model likelihood; see Burnham & Anderson 2002). Non-nested models within 2 $AIC_C$ of each other were considered equally well supported. Confidence intervals were obtained by nonparametric bootstrapping of the model parameters using 10 000 replicates.

2.3. Population viability analyses

Population viability analyses (PVA) were carried out separately for each colony. In each case, 10 000 future population trajectories were modelled using Eq. (1). The quasi-extinction threshold was set at 20 pairs. Confidence limits around the median population trajectory were estimated as population predic-
A population prediction interval is ‘the stochastic interval that includes the unknown population size at a specified future time with a given probability or confidence level’ (Lande et al. 2003, p. 108) and incorporates stochasticity as well as parameter uncertainty. The effects of demographic and environmental stochasticity are included via Eq. (1). Uncertainty about parameter estimates was taken into account by simulating the population time series using the estimated parameters 10 000 times and re-estimating the parameters from each simulation. This method produces sampling distributions for all parameters ($\bar{r}$, $\sigma^2$, and $\beta$), from which a random set of population parameters is drawn, and accounts for the presence of sampling correlation (Lande et al. 2003). No PVA was performed for Sklinna because the population is already extinct and crossed the quasi-extinction threshold of 20 pairs for the first time in 2001.

For each colony, a set of at least 3 different PVAs was carried out. The first PVA was based on the null population model, i.e. without covariates. The remaining PVAs were based on the estimates derived from the best population model(s) incorporating SST as a covariate; half of the latter PVAs assumed average SST to stay constant at the actual level of the years 2000 to 2011, the other half assumed average SST to increase in line with predictions of ocean warming (see Fig. 3). Using this approach, it is possible to directly compare the viability of each colony under different assumptions (effect of SST present vs. absent, and warming present vs. absent).

### 2.4. Climatic variables

The climatic covariate considered as an explanatory variable was SST, based on the Extended Reconstruction SST data set available on a $2^\circ \times 2^\circ$ grid (ERSST v 3b, NOAA 2012; cf. Smith et al. 2008). We considered spring and summer SSTs around each of the breeding colonies calculated as seasonal means (March to May and June to August, respectively) of the 2 grid cells adjacent to the colonies (Runde, 62−64° N, 2−8° E; Sklinna, 64−66° N, 8−14° E; Vedøy, 66−68° N, 8−14° E; Hjelmsøya, 70−72° N, 22−28° E; Hornøya, 70−72° N, 28−34° E). SSTs for the non-breeding season were taken from the following areas and periods: autumn SST off Svalbard was defined as the spatial mean September SST within the area 74−80° N, 14−36° E; winter SST in the Grand Banks area as the spatial and seasonal mean SST during November to January within the area 40−62° N, 38−60° W. These choices of areas and periods were based on the actual spatiotemporal distribution of kittiwakes from the relevant colonies outside the breeding season (Frederiksen et al. 2012).

SSTs were considered as covariates at different time lags, allowing for different biological explanations of
potential effects: if SST affects breeding propensity (i.e. absence/presence of adult birds during the population count), this would be visible in population models as an unlagged effect of SST. In contrast, if SST affects adult survival after breeding, the corresponding change in population size would not become evident before the population count of the following year, showing up in the population model as an SST effect lagged by 1 yr. Effects of SST on recruitment would entail even longer time lags: most kittiwakes that return to their breeding colony to breed do so at an age of 3 to 4 yr (Coulson 2011). If breeding success (or survival of juveniles during their first winter) is affected by SST, the corresponding change in population size would therefore not be counted before the cohort affected recruits to the breeding population, i.e. 3 to 4 yr later. In a population model, an SST effect on reproduction would thus become evident as an effect of SST lagged by the number of years that corresponds to the mean age at recruitment. Based on these assumptions, we considered SSTs around colonies at time lags of 0, 1, 3 and 4 yr; SSTs off Svalbard and in the Grand Banks area were considered at time lags of 1, 3 and 4 yr (where the time period from autumn or winter to the following breeding season is considered to be a time lag of 1).

Estimates of SST in a future climate scenario were extracted from the Norwegian Earth System Model (NorESM; Iversen et al. 2013), a global coupled climate model. The scenario chosen was RCP8.5 (Representative Concentration Pathway) with radiative forcing target level at 8.5 W m⁻² in 2100, which is a very high baseline emission scenario leading to CO₂ concentrations at 1370 ppm in 2100 (van Vuuren et al. 2011). The RCP8.5 scenario does not include any specific climate mitigation target, corresponding to a doubling in greenhouse gas emissions by 2050 and 3-fold increase by 2100 (Riahi et al. 2011). SSTs were extracted for the exact positions of each colony and for points within the wintering areas (off Svalbard, 78° N, 26° E; Grand Banks, 51° N, 46° W).

For each of these points, SSTs for the period 2006 to 2100 were used to estimate a linear trend. The period of overlap between the ERSST and NorESM data (2006 to 2011) was used to adjust the historical with the future time series to ensure that the projected SST trend started at the same value as the empirical SST data end point. For each of the 10 000 PVA runs, an independent SST time series was generated. This time series consisted of white noise (zero mean and assuming the same variance as in the past, inferred from the relevant ERSST time series), added either to the mean SST of the years 2000 to 2011 or to the projected SST trend (Fig. 3).

All models were run in the R environment (R Development Core Team 2011). Estimates are provided with 95% confidence intervals.

3. RESULTS

The 5 Norwegian kittiwake Rissa tridactyla colonies studied (Runde, Sklinna, Vedøy, Hjelmsøya and Hornøya) declined steeply during the study period (Fig. 2). Beyond the decline, there was no strong temporal covariation among the colonies. If the negative trend was not removed, population counts were highly correlated (except Sklinna, all pairwise R > 0.7, all p < 0.01; Sklinna was only correlated to Vedøy). Upon removal of the trend, however, counts were uncorrelated (all |R| < 0.3, all p > 0.18). Nor were annual population growth rates correlated across colonies (whether de-trended or not, all |R| < 0.4, all p > 0.14). Population synchrony across colonies was thus virtually absent.

The long-term mean rate of decline was similar in all colonies; however, the temporal variability was much higher in some colonies (Fig. 2). This is evident from the best population model without covariates (Table 1), which assumed a common long-term intrinsic population growth rate $\bar{r}$ of $-0.055 \pm 0.026$
Table 1. Population models for 5 Norwegian black-legged kittiwake populations, fitted to all populations simultaneously and assuming Brownian population dynamics. Covariates used were sea surface temperatures from different ocean areas. Models are sorted by decreasing support and presented using estimates and 95% confidence intervals (CI), number of parameters (K), ΔAIC_C and model likelihood (ML). **Bold**: best-supported models. Models with covariates assumed a common growth rate (\(\bar{r}\)) and separate environmental variances (\(\sigma^2_e\)) in the 5 colonies (which was the best supported parameterisation of models without covariates). The 2 top models had AIC_C weights of 0.40 and 0.38, respectively. See Table S1 for density-dependent models.

<table>
<thead>
<tr>
<th>Model: covariate (time lag, yr)</th>
<th>Estimate</th>
<th>CI</th>
<th>K</th>
<th>ΔAIC_C</th>
<th>ML</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grand Banks (3)</td>
<td>(b = -0.079)</td>
<td>(-0.132) to (-0.027)</td>
<td>7</td>
<td>0.00</td>
<td>1.00</td>
</tr>
<tr>
<td>Svalbard (1)</td>
<td>(b = -0.244)</td>
<td>(-0.404) to (-0.081)</td>
<td>7</td>
<td>0.12</td>
<td>0.942</td>
</tr>
<tr>
<td>Svalbard (1), de-trended</td>
<td>(b = -0.240)</td>
<td>(-0.450) to (-0.035)</td>
<td>7</td>
<td>3.60</td>
<td>0.165</td>
</tr>
<tr>
<td>Grand Banks (1)</td>
<td>(b = -0.059)</td>
<td>(-0.113) to (-0.004)</td>
<td>7</td>
<td>4.07</td>
<td>0.131</td>
</tr>
<tr>
<td>Grand Banks (3), de-trended</td>
<td>(b = -0.088)</td>
<td>(-0.174) to (-0.004)</td>
<td>7</td>
<td>4.40</td>
<td>0.111</td>
</tr>
<tr>
<td>Colonies (1)</td>
<td>(b = -0.065)</td>
<td>(-0.133) to +0.003</td>
<td>7</td>
<td>5.12</td>
<td>0.077</td>
</tr>
<tr>
<td>No covariate, common (\bar{r}), separate (\sigma^2_e)</td>
<td>(\bar{r} = -0.055)</td>
<td>(-0.081) to (-0.030)</td>
<td>6</td>
<td>6.42</td>
<td>0.040</td>
</tr>
<tr>
<td>Grand Banks (1), de-trended</td>
<td>(b = -0.031)</td>
<td>(-0.116) to +0.052</td>
<td>7</td>
<td>8.09</td>
<td>0.018</td>
</tr>
<tr>
<td>Colonies (1), de-trended</td>
<td>(b = -0.027)</td>
<td>(-0.142) to +0.085</td>
<td>7</td>
<td>8.44</td>
<td>0.015</td>
</tr>
<tr>
<td>No covariate, separate (\bar{r}), separate (\sigma^2_e)</td>
<td>(\bar{r} = -0.070)</td>
<td>(-0.106) to (-0.035)</td>
<td>2</td>
<td>45.64</td>
<td>0.000</td>
</tr>
<tr>
<td>No covariate, common (\bar{r}), common (\sigma^2_e)</td>
<td>(\bar{r} = -0.027)</td>
<td>(-0.142) to +0.085</td>
<td>7</td>
<td>8.44</td>
<td>0.015</td>
</tr>
</tbody>
</table>

(corresponding to an annual reduction of 5.7 ± 2.7%), while the environmental variance \(\sigma^2_e\) differed tenfold between 0.011 ± 0.006 at Vedøy and 0.118 ± 0.078 at Sklinna. Variability was not merely a function of colony size; for instance, Hjelmsøya was a larger and more variable colony than Hornøya (correlation between mean log population size and environmental variance, \(R = -0.83, p = 0.080, n = 5\)).

The best model without covariates could be improved using SST as covariate (Table 1). In all cases, SST was negatively related to population growth rate. The 2 best-supported models indicated that SST during the non-breeding season had the strongest effect. According to the first model, population growth rate declined 3 yr after a warm winter in the Grand Banks area. This time lag suggests an effect on the first-year survival of future recruits. The variance in population growth rate explained by this effect was 14% in Runde, 11% in Hornøya, 6% in Vedøy and <1% in Sklinna and Hjelmsøya. According to the second model, population growth rate declined in years following a warm autumn southeast of Svalbard. This unlagged effect suggests an effect on adult return rate and explained a fifth of the variance in population growth rate in the 2 northernmost colonies (21% in Hornøya, 19% in Hjelmsøya, <5% elsewhere). When de-trending these 2 covariates, the corresponding models were somewhat poorer but still at least 2 AIC_C units better than models without covariates (Table 1). This indicates that the model support is not merely due to unrelated trends in population growth rate and temperature.

Of the remaining covariate models, 2 were better supported than the model without covariates, but somewhat poorer (ΔAIC_C > 2) than the 2 top-ranked models. These indicated an effect of SST at the Grand Banks in the previous winter and of SST around the colonies in the previous summer (Table 1). No combinations of 2 or more covariates achieved more support than single-covariate models.

When fitting separate models to each colony, results were somewhat different (Table 2). The best models for Hornøya and Hjelmsøya contained SST southeast of Svalbard in the previous autumn. In the case of Hjelmsøya, this model could be improved by adding the previous year’s SST around the colony. The latter effect was estimated to be positive after SST southeast of Svalbard was accounted for. The effect of winter SST in the Grand Banks area 3 yr earlier was only supported as a covariate to the population dynamics at Runde. The population dynamics at Sklinna and Vedøy could not be explained using the covariates considered (Table 2; cf. Table S2 in the Supplement).

Based on models without covariates, all extant colonies except Hornøya had a median time to extinction of <90 yr and a lower 95% confidence limit of <50 yr (Table 3). Estimated extinction probabilities increased when using covariate models and especially when adding a warming trend to the predicted future SST values. This pattern is evident from Fig. 4, which shows population trajectories for 1 colony under the 3 different assumptions. For all colonies, median time to extinction was significantly shorter.
under the assumption of a causal link to SST and a warming trend (reduced by as much as 56%, 46%, 81% and 90%, respectively) than without these assumptions (Table 3, Fig. 4).

One of the models (Hjelmsøya) had covariates with opposite signs (Table 2). While this lead to increased estimates of time to extinction compared to the 1-parameter model under the assumption of constant SST, the 2- and 1-parameter models did not differ under the assumption of warming (Table 3).

4. DISCUSSION

4.1. Retrospective models of population dynamics

Our analyses of population dynamics of 5 Norwegian kittiwake *Rissa tridactyla* colonies have shown that their overall rates of population decline were similar (~5.7% yr⁻¹), although we found no evidence of synchrony among their annual changes in breeding numbers. SSTs in different areas of the North Atlantic explained between 6 and 37% of the inter-annual variation in population growth rates, but regions and time lags differed between the colonies. Most slopes were estimated to be negative, i.e. warmer conditions were related to stronger population decrease. We ascertained that the effects of SST were not just artefacts created by uncorrelated trends by verifying the findings with de-trended time series.

The oceanic regions considered had been chosen based on recent evidence of the non-breeding distribution of kittiwakes from these colonies, and the regions identified as most relevant by the population models are fully compatible with this evidence. Geolocator data singled out 2 areas as especially important for adult kittiwakes outside the breeding season (Frederiksen et al. 2012, B. Moe et al. unpubl. data): an area southeast of Svalbard, visited by kittiwakes from the northernmost colonies after the breeding season (September), and the Grand Banks, visited by birds from all colonies in winter (November to January). This explains why, in our analyses, SST east of Svalbard in September accounts for roughly 20% of population dynamics at Hornøya and Hjelmsøya, which are the colonies utilising this area the most. Models incorporating SST at the Grand Banks were better than the null model and/or the best supported covariate model for all colonies except Hjelmsøya, although this variable entered the optimal model for Runde only, where it explained some 12% of the inter-annual variation in population dynamics. Unfortunately, no geolocator data are available from Runde and Sklinna. However, based on our findings, it is unlikely that these colonies deviate from the multi-colony pattern revealed by geolocators in other colonies.

The absence of population synchrony between the colonies is somewhat surprising given the evidence that birds from several breeding colonies use the same oceanic regions during winter. This is an important finding in itself, as the degree of population synchrony may affect the extinction probability (Engen et al. 2002). The most likely explanation for the absence of synchrony is the presence of environmental noise and measurement error.
Environmental noise may not only explain the absence of population synchrony but also be invoked as an alternative explanation for the population decline as such. Factors that have been documented to affect population dynamics in other areas and/or other species of seabirds include predation pressure (e.g. by white-tailed eagles Haliaeetus albicilla, Hipfner et al. 2012), competition with larger gulls (e.g. Oro et al. 2009) or interactions with commercial fisheries (e.g. Frederiksen et al. 2004). While such factors may have also contributed to the decline in some Norwegian colonies, they cannot explain the overall pattern. The population decline has a very similar slope in colonies, they cannot explain the overall pattern. The column ‘50%’ provides the median time to extinction, and the column ‘2.5%’ provides its lower 95% confidence limit. *Times to extinction that differ significantly from a model without covariates and warming (i.e. a median time to extinction of less than the 2.5% quartile of the corresponding null model). Bold: threshold values that correspond to Red List criteria (50% within 10 yr: critically endangered; 20% within 20 yr: endangered; 10% within 100 yr: vulnerable)

<table>
<thead>
<tr>
<th>Model: covariate (time lag, yr)</th>
<th>Time to extinction (yr) with a probability of</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>50%</td>
</tr>
<tr>
<td><strong>Runde</strong></td>
<td></td>
</tr>
<tr>
<td>Null</td>
<td>79</td>
</tr>
<tr>
<td>Grand Banks (3)</td>
<td>45</td>
</tr>
<tr>
<td>Grand Banks (3) + warming</td>
<td>35*</td>
</tr>
<tr>
<td><strong>Vedøy</strong></td>
<td></td>
</tr>
<tr>
<td>Null</td>
<td>89</td>
</tr>
<tr>
<td>Grand Banks (3)</td>
<td>60</td>
</tr>
<tr>
<td>Grand Banks (3) + warming</td>
<td>48*</td>
</tr>
<tr>
<td><strong>Hjelmsøya</strong></td>
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<tr>
<td>Null</td>
<td>52</td>
</tr>
<tr>
<td>Svalbard (1)</td>
<td>35</td>
</tr>
<tr>
<td>Svalbard (1) + warming</td>
<td>10*</td>
</tr>
<tr>
<td>Svalbard (1) + colony (1)</td>
<td>49</td>
</tr>
<tr>
<td>Svalbard (1) + colony (1) + warming</td>
<td>9*</td>
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<td><strong>Hornøya</strong></td>
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<td>Null</td>
<td>181</td>
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<tr>
<td>Svalbard (1)</td>
<td>94</td>
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<tr>
<td>Svalbard (1) + warming</td>
<td>18*</td>
</tr>
</tbody>
</table>

SST at the Grand Banks was mostly relevant if lagged by 3 yr (Runde, Vedøy and Hornøya). A time lag of 3 yr is compatible with an effect of SST on recruitment, i.e. on the survival of immature birds (cf. Sandvik et al. 2012). Fledglings that later recruit to their natal colony will enter the population count when they first return to the breeding colony to build a nest (not necessarily to lay eggs), which in kittiwakes happens at 3 to 4 yr of age (Coulson 2011). The findings thus suggest that recruitment is poor in cohorts that experience warm conditions during their first winter after fledging. This interpretation does not involve the assumption that immatures were unaffected by environmental conditions in other years; however, consistently finding a 3 yr lag across cohorts is convincing evidence that the effect was strong enough not to be masked by the environmental conditions of the intervening seasons. On the other hand, the interpretation presupposes that immatures use the same areas as adults. Geolocator data from immatures that would allow us to test this assumption are, however, not available at present. Time lags in climate ecology can also originate from effects that are mediated through...
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Although this alternative explanation is less likely in this case, because kittiwakes seem to feed at a low trophic level during winter, it cannot currently be ruled out entirely.

The findings thus corroborate results from the North Sea, where adult survival was negatively related to SST (Frederiksen et al. 2004, 2006); in contrast, the present study did not find effects of SST on chick production (which would have resulted in a 3 yr lag of local SST), but rather on first-year survival.

A study of adult survival rates of kittiwakes from the Hornøya colony can shed some light on the likely mechanisms underlying population responses to SST. Adult survival was strongly affected by prey abundance, notably of capelin in the Barents Sea and of sea butterflies (Thecosomata) at Grand Banks (Reiertsen et al. in press). Capelin and sea butterflies are known to be important prey of kittiwakes during summer and autumn, respectively (Barrett 2007, Karlovsky et al. 2008), and may represent the causal link between SST and survival. No biotic link explaining the importance of the region off Svalbard has been identified so far (Reiertsen et al. in press).

Local conditions, i.e. summer SST around the colonies, entered the population model of Hjelmsøya but only after the effect of the non-breeding season had been taken into account (Table 2). This is the only covariate estimated to have a positive slope, indicating that adult survival was higher after warmer breeding seasons.

4.2. Population viability analyses

The kittiwake is currently classified as endangered in the Norwegian Red List (Kålås et al. 2010). This decision was based on Criterion A2b, because the Norwegian mainland population has decreased by almost 80% within 3 generations. According to the PVAs of the present study, most Norwegian colonies would be categorised as vulnerable (10% extinction risk within 100 yr; see Table 3) when applying Red
List criterion E (IUCN 2001) to each of them. In the absence of any covariate effect, Hornøy is the most viable colony, classified as near threatened (5% extinction risk within 100 yr), which would change to critically endangered (50% extinction risk within 10 yr) according to the PVA that includes a warming trend. The Hjelmsøy colony even crosses the threshold to critically endangered (50% extinction risk within 10 yr) according to PVAs that include warming trends (Table 3). The Sklinna population, which was the smallest of the colonies studied, went extinct during the study period.

These results are only indicative because Red List criteria are not applicable to single populations (Hartley & Kunin 2003). For example, local populations may have a high turnover rate, without the species as such being at threat (viz., if new populations are established at the same rate as other populations go extinct). There is no indication that this is the case for kittiwakes. Moreover, the negative trend was very similar in all study colonies, which covered a large part of the species’ breeding range in Norway. No comparable time series are available from other Norwegian colonies, but there is no indication that other colonies are better off than the 5 colonies studied here (Erikstad & Systad 2009).

The warming scenario may have overestimated the extinction risk (or underestimated time to extinction) for several reasons. First, the warming model chosen corresponds to a ‘business as usual’ scenario, which may be too pessimistic an assumption. Our main purpose was to compare 2 extreme models, one assuming constant SST and the other a drastic but realistic warming trend. As such, these 2 models represent reasonable limits that embrace the actual future trend. Most emission scenarios do, however, assume trends that are closer to the ‘business as usual’ scenario than to a ‘no change’ scenario (van Vuuren et al. 2011).

Second, the PVAs assume that the mechanisms of the past remain unchanged in the future, which, of course, is uncertain. For example, we do not yet know a great deal about the temporal stability of foraging patterns and wintering areas of kittiwakes. If they remain stable, the PVAs offer realistic viability estimates. However, at least 2 factors may decouple kittiwake population dynamics from autumn/winter SST: (1) kittiwakes may follow their main prey species as they migrate to other (presumably colder) areas, and/or (2) kittiwakes may shift to other prey. The latter may occur either because the current prey species are replaced by more warm-tolerant species originating from more southerly waters or because kittiwakes move to areas where more warm-tolerant species are abundant during winter.

A third factor that might have resulted in overestimated extinction risks is that past (actual) and future (modelled) SSTs may not be directly comparable. SST is not sampled to the same extent in all areas. A coarser sampling in Arctic waters would, for example, cause underestimation of inter-annual SST variation in the ERSST dataset. This would, in turn, result in inflated slopes in our population models, which would then overestimate the effect of warming. Further studies are needed to rule out this potential source of systematic errors.

Even when excluding a warming trend, however, the PVAs with SST covariates suggested 33 to 48% shorter median times to extinction than PVAs without covariates (Table 3, Fig. 4). Because of the high uncertainty and correspondingly wide population prediction intervals, these reductions are not statistically significant, although they certainly would be biologically so. The reasons for these drastically increased extinction risks are the negative effect of SST and the fact that even constant SST at current levels represents conditions that are considerably warmer than the long-term average (cf. Fig. 3).

Frederiksen et al. (2012) have hypothesised that the decline of kittiwakes in the North Atlantic may be due to environmental conditions at the Grand Banks, which is an overwintering area that seems to be common to the whole Atlantic population. Our findings from Runde, and to some degree from Sklinna and Vedøy, are compatible with this hypothesis. However, the decline in the 2 northernmost colonies (Hjelmsøy and Hornøy) was more closely related to autumn conditions off Svalbard, while the population declines were similar. The support for the importance of the Grand Banks area is thus somewhat equivocal.

In conclusion, although ocean warming is not the sole explanation for the decline of Norwegian kittiwake populations, it aggravates the situation considerably. Unless kittiwakes are able to switch to other foraging areas or prey, especially outside the breeding season, the populations surveyed will reach quasi-extinction within a couple of decades.

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