INTRODUCTION

There is now very strong evidence for a progressive effect of human activity on global climates, with increases in atmospheric CO₂ levels strongly suspected to have induced increases in air temperatures, particularly over the past 50 yr (IPCC 2007). Less certain, however, are the long-term impacts of rising temperatures on different species and ecosystems (Hockey & Midgley 2009, Grémillet & Boulinier 2009). In marine environments, warming of surface waters results in a low-density surface layer that greatly reduces vertical mixing, with a consequent reduction in nutrient inputs into the euphotic zone and hence in the primary productivity of surface waters during the summer period (Richardson & Schoeman 2004, Behrenfeld et al. 2006). In polar regions, this effect may then be exacerbated by decreases in surface-water salinity due to increased fresh-water input as a result of the melting of coastal and inland ice as well as Arctic permafrost (Jacobs et al. 2002, Greene & Pershing 2007). However, surface warming can also directly favour phytoplankton growth at high latitudes (Behrenfeld & Falkowski 1997), and in all regions, rising air temperatures can generate large atmospheric pressure differentials, resulting in high wind stress on the surface of the oceans. This in turn increases the upwards transfer of nutrients and so enhances primary productivity (Toggweiler &
Russell 2008). The net effect of these antagonistic processes varies regionally but global ocean primary productivity, as assessed by remote sensing, has decreased significantly since 1999 (Fig. 1 in Behrenfeld et al. 2006). This decrease is highly correlated with large-scale climatic indices, strongly suggesting that global warming has had a net negative impact on marine primary productivity (Le Bohec et al. 2008, Grémillet & Boulinier 2009).

There is growing evidence that decreases in primary productivity can be propagated up the food chain (Beaugrand et al. 2002, Behrenfeld et al. 2006), extending in some cases to higher marine predators such as piscivorous seabirds (e.g. Gjerdrum et al. 2003, Frederiksen et al. 2006). Rising temperatures may also have direct physiological effects on both predatory fish (Perry et al. 2005) and seabirds (Oswald et al. 2008), in some cases exacerbating the impacts of changes in primary productivity (Hamer 2010, Oswald et al. in press). Intense fishing pressure on predatory fish can also have cascading effects on marine food webs (Watermeyer et al. 2008, Baum & Worm 2009) and the greatest threat to fish stocks is likely to be the combined effects of climate change and overfishing (Brander 2007). Recent studies have suggested that these combined effects can also have important consequences for seabird breeding success, survival and population stability (Frederiksen et al. 2004b, Ainley & Blight 2009).

In the Northeast Atlantic Ocean, several species of seabird at sites bordering the North Sea have experienced delayed breeding, reduced breeding success and, in some cases, declines in population sizes since the 1990s (Frederiksen et al. 2004a, Mavor et al. 2006). These events have been linked to reductions in the abundance and nutritional quality of their main prey (lesser sandeels Ammodytes marinus), which were in turn associated with warming of surface waters (Frederiksen et al. 2004b, Wanless et al. 2007). In particular, an inflow of warm nutrient-poor oceanic water, together with stronger stratification of surface waters, has favoured a northward expansion of warm-water copepod species and a decline in cold-water zooplankton (Beaugrand 2004, Grémillet & Boulinier 2009). Together with impacts of industrial sandeel fisheries (Frederiksen et al. 2004b, Poloczanska et al. 2004), these changes in plankton assemblages have resulted in poor and erratic growth and survival of sandeels, with adverse consequences for North Sea seabirds (Frederiksen et al. 2007). However, studies elsewhere in the region have not previously recorded such adverse effects (Swann 2000, Votier et al. 2005), despite the fact that sea surface temperature has also been rising in these areas (ICES 2007). Indeed, studies at more northerly latitudes have recorded or suggested positive relationships between sea surface temperature and vital rates (breeding success, recruitment, adult survival, etc.) of several species (Aebischer et al. 1990, Hamer et al. 1991, Thomson & Ollason 2001, Durant et al. 2003, D’Alba et al. 2010). Moreover in Shetland, recent low abundance of sandeels coincided with record-high abundance of herring Clupea harengus (Frederiksen et al. 2007), in keeping with the notion that food webs in cooler waters may be more strongly influenced by ‘top-down’ effects of predation than by ‘bottom-up’ effects of changes in primary and secondary production (Hunt et al. 2002, Frank et al. 2007).

Seabirds differ in their vulnerability to adverse effects of low prey availability (Furness & Tasker 2000). For instance, species foraging relatively close to the colony are typically more vulnerable than longer distance foragers, because they have less flexibility in their foraging locations and time-activity budgets (Diamond 1978, Hamer et al. 2007). Procellariiform seabirds (albatrosses, petrels, shearwaters, etc.) typically forage at great distances from the colony (Brooke 2004), and several species have been recorded to employ a dual-foraging strategy entailing a combination of short foraging trips mainly used to feed the chick, interspersed with longer trips during which the adult rebuilds its reserves (Weimerskirch et al. 1994, Weimerskirch 1998). There is also growing evidence that procellariiforms can adjust their food provisioning strategy according to food availability and foraging costs (Granadeiro et al. 1999, Duriez et al. 2000, Quillfeldt et al. 2007). For these reasons, procellariiforms have been viewed as the least vulnerable to changes in the distribution and abundance of their prey (Furness & Tasker 2000).

Manx shearwaters Puffinus puffinus breed in large colonies around the Northeast Atlantic (Brooke 1990). They typically make foraging trips lasting 1 to 3 d with little indication of a dual-foraging strategy (Gray & Hamer 2001, Hamer et al. 2006). On the islands of Skomer and Skokholm, Southwest Wales, they have been studied sporadically since the 1920s and annually since about 1990, yet there have been no accounts of feeding conditions that have led to poor chick growth or breeding failures (Lockley 1930, Harris 1966, Brooke 1990, Hamer & Hill 1997). During chick-rearing, Manx shearwaters from these islands prey mainly on clupeid fish such as herring Clupea harengus and sprat Sprattus sprattus (Brooke 1990), with foraging effort concentrated in the Irish and Celtic Seas (Guilford et al. 2008). Stocks of herring in the Celtic Sea have declined steadily following a recovery in the 1980s and have reached critically low levels in recent years (ICES 2007). There are no comparable data for stock sizes of sprat in the region or herring in the Irish Sea, but they are also thought to be low (ICES 2007). Here, we use data for years spanning the period from 1965 to 2008 to investigate annual variation in the phe-
nology, breeding success and foraging behaviour of Manx shearwaters in Southwest Wales and to relate these data to annual variation in climatic and prey population indices.

**MATERIALS AND METHODS**

Fieldwork was carried out at the large colony of Manx shearwaters (>100 000 pairs) on Skomer Island, Southwest Wales (51.73° N, 5.28° W), each year from 1995 to 2008 and on the neighbouring island of Skokholm (ca. 4 km away) from 1965 to 1967 and from 1973 to 1976. Manx shearwaters are burrow nesters, and chicks were reached via short access tunnels in the roof of each burrow, capped with removable earth lids. This system reduced overall disturbance and had no adverse effects on burrow occupancy, food-provisioning rates or breeding success (Brooke 1990, Hamer & Hill 1997). We determined hatching dates of chicks (between 40 and 60 nests each year, with mostly different study burrows being sampled each year) by direct observation or by calibrating wing length (maximum flattened chord measured to the nearest 1 mm) against wing growth in chicks of known age (Brooke 1990). Chicks were then weighed daily to the nearest 5 g using a Salter spring balance (RS Components). Manx shearwater chicks gain weight for about 50 to 55 d post-hatching before losing mass prior to fledging at around 70 d. Mass growth was monitored until fledging in 1965, 1973–1976, 1995, 1998–2000 and 2007–2008; peak and fledging masses are described for these years only. In addition, every night from the end of August in 2007 and 2008, we caught and weighed a sample group of young Manx shearwaters on the surface prior to fledging for comparison with published Skokholm data from 1965 to 1967 (Perrins et al. 1973).

**Food provisioning behaviour of adults.** Parents are nocturnal at the colony, and overnight nest attendance was monitored by simultaneously radio-tracking both adults at each of 10 nests in 1999 and 2002, 21 nests in 2007 and 19 nests in 2008. In each case, both adults were caught when chicks were 15 to 25 d old, weighed and fitted with a VHF radio-transmitter (Biotrack) weighing 2 g in 1999 and 2002 and 4.5 g in 2007 and 2008 (<1.5% of adult body mass). Transmitters were attached to the central tail feathers using self-amalgamating tape (RS Components). Attendance of each adult at the colony was then monitored using a scanning receiver for an average of 15 and 16 consecutive nights in 1999 and 2002, respectively, and until chicks reached ca. 60 d of age in 2007 and 2008. Trials at the colony indicated that weak and intermittent signals were sometimes received if a bird was on the water in the vicinity of land, but consistent, strong signals were obtained only from birds returning to their nests (Riou & Hamer 2008). Chicks were weighed twice daily (at dawn and dusk), and mass increments between successive weighings overnight were used as estimates of meal sizes.

There was no discernible effect of radio-tracking on chick growth in 1999 or 2002 (Gray & Hamer 2001, Quillfeldt et al. 2004). As a further check in 2007 and 2008, we compared the growth of chicks at nests where both parents had been radio-tracked with a separate sample of 30 nests where parents were never handled. There was no significant effect of radio-tracking on average chick peak mass (± SE) (radio-tracked: 505.1 ± 11.9 g; controls: 515.7 ± 12.6 g; likelihood ratio test: $\chi^2_1 = 0.1$, $p = 0.7$), nor was there an effect on fledging mass (radio-tracked: 394.5 ± 11.4 g; controls: 393.3 ± 7.2 g; $\chi^2_1 = 0.01$, $p = 0.9$).

**Environmental variables.** Monthly sea surface temperature (SST) data were obtained from the UK Meteorological Office (Hadley Centre; HadISST 1.1 Global sea-ice coverage and SST 1870 to present, available from http://climexp.knmi.nl) and used to calculate the average winter (December to April) SST for waters surrounding Skomer. Winter North Atlantic Oscillation (NAO) index values were obtained from the Climatic Research Unit, Norwich, UK. We used data for these months to characterize climate over the period preceding the birds’ breeding season each year. Both winter SST and winter NAO have previously been linked to fish and seabird productivity (Borges et al. 2003, Frederiksen et al. 2004b), and in the Baltic Sea, mild winters (as indicated by a positive winter NAO index) have a positive effect on year-class strength of herring in spring (Axenrot & Hansson 2003). In the waters around Skomer, herring spawn on the south coast of Ireland during the winter and move to feeding grounds in the Celtic and Irish Sea during the summer. Here, we used spawning stock biomass (SSB; data from the ICES Her-Ring Working Group; ICES 2009) as an index of prey abundance each year. The modal length of prey regurgitated by birds at the colony (15 cm; Brooke 1990) best corresponds to fish aged 2 yr, and so we used SSB 2 yr previously, which accurately predicts year-class strength of age-2 herring (Axenrot & Hansson 2003). We also examined prey quality, in terms of the mean mass of individuals aged 2 yr (data from ICES 2009).

**Statistical analyses.** Chick growth and food provisioning were compared across years using general linear modelling in R (www.R-project.org). Hatching date and chick peak mass were first analysed as a function of year using $F$-tests. Growth rates during the phase of linear growth (10 to 30 d post-hatching) were examined by modelling chick mass as a function of year and age, with chick ID included as a random effect to account for repeated measurements (Pinheiro & Bates 2000).
Hatching date and peak mass were then analysed as a function of SST, NAO and herring mass and abundance, starting with all the variables in the initial model and including year as a random effect. Finally, parental attendance was modelled as a function of year in a binomial model with parent ID included as a random effect. Attendance was measured as the probability of an individual bird being present at the colony each night during the period with the most intense and least variable food provisioning rhythm when chicks are between 20 and 50 d of age (Hamer & Hill 1997). In all cases, model simplification was carried out by sequential removal of the least significant terms, using likelihood ratio tests ($F$-tests and, when a random effect was included, $\chi^2$ tests). All significant terms remained in the minimum model used for statistical inference. All models met the assumptions of normality and homoscedasticity.

RESULTS

Timing of breeding and growth of chicks

Hatching occurred significantly later in both 2007 and 2008 than in previous years ($F_{16, 734} = 3.3$, $p < 0.0001$; Fig. 1), but there was no annual trend in hatching date ($\chi^2_1 = 1.4$, $p > 0.2$) and no effect of either NAO or winter SST (all $\chi^2_1 < 1.2$, $p > 0.3$). Chick mass during the linear phase of growth differed significantly between years ($\chi^2_3 = 225$, $p < 0.0001$), with an average growth rate of $14.9 \pm 0.1$ g d$^{-1}$ from 1965 to 1976, $12.7 \pm 0.2$ g d$^{-1}$ from 1995 to 2000 and $11.3 \pm 0.3$ g d$^{-1}$ in 2007 (Fig. 2). In 2008, chicks had an intermediate rate of growth to 30 d post-hatching ($13.2 \pm 0.3$ g d$^{-1}$), but in both 2007 and 2008, chicks showed very poor growth from then until fledging (Fig. 2).

Peak masses of chicks decreased significantly with year ($F_{1, 411} = 225$, $p < 0.0001$) and were lower in 2007 and 2008 than in all previous years (Fig. 2). Variation in peak mass closely matched SST in the immediately preceding winter in the waters around the colony ($\chi^2_1 = 10.6$, $p = 0.001$; Fig. 3) but was not related to winter NAO (all $\chi^2_1 = 0.5$, $p = 0.5$). In a separate model dealing with the effects of herring mass and abundance, chick peak mass was strongly dependent on mean body mass of 2 yr old herring ($\chi^2_1 = 8.6$, $p = 0.003$; Fig. 4) but was not significantly related to spawning stock biomass ($\chi^2_1 = 2.8$, $p = 0.09$). In a model that included both SST in the preceding winter and herring body mass, both terms showed significant independent effects on chick peak mass ($\chi^2_1 = 4.5$, $p = 0.03$ and $\chi^2_1 = 5.4$, $p = 0.02$, respectively).

Body masses of fledglings were substantially lower in 2007 and 2008 than from 1965 to 1967 (Fig. 5). Chicks that fledged at earlier dates in 2008 were heavier than later fledglings that year and than all fledglings in 2007 (interaction between date and year; $F_{1, 380} = 10.7$, $p = 0.001$; Fig. 5).

Parental attendance and food provisioning

The probability of a parent returning to the nest overnight differed significantly between years ($\chi^2_3 = 20.7$, $p = 0.0001$), being substantially lower in 2007 and
than in 1999 and 2002 (Table 1). Adults increased the frequency of long trips (>3 d) and reduced the frequency of 2 to 3 d trips in 2007 and 2008 compared to 1999 and 2002 ($\chi^2_1$ on a $2 \times 2$ contingency table) = 25.0, $p < 0.001$; Fig. 6). The average duration of long trips also increased substantially, from 4.8 ± 0.4 d (maximum = 7 d) in 1999 and 2002 to 5.6 ± 0.1 d (maximum = 10 d) in 2007 and 2008. These changes resulted in a bimodal distribution of trip durations in 2007 and 2008, which was not present in 1999 or 2002 (Fig. 6).

The sizes of individual meals delivered by parents showed little variation among years ($\chi^2_3 = 0.7$, $p = 0.9$; Table 1). However, double meals, resulting from delivery of food by both parents, showed significant variation among years ($\chi^2_3 = 36.8$, $p < 0.0001$), being heavier in 2007 and 2008 than in earlier years. They were close to twice the mass of single meals only in 2007 and 2008.

**DISCUSSION**

In 2007 and 2008, Manx shearwaters in SW Wales bred later and chicks reached substantially lower peak masses than in any of the years since the 1960s for which
The observed decrease in adult attendance was the result of a change in foraging behaviour whereby parents increased both the duration and frequency of long trips, with a decrease in the frequency of 2 and 3 d trips. Parents thus switched from using mainly short foraging trips in 1999 and 2002 to a dual-foraging strategy in 2007 and 2008. This change in foraging behaviour is similar to that seen in Cory’s shearwater Calonectris diomedea, which also switched to dual-foraging under conditions of low prey availability (Granadeiro et al. 1998, Magalhaes et al. 2008). The sizes of single meals were not affected in this study, but the average yield to chicks from single meals was much lower in 2007 and 2008 than in 1999 and 2002 (21.7 g d⁻¹ and 27.8 g d⁻¹, respectively; calculated from data in Table 1) as a result of longer intervals between feeds. Double meals were substantially smaller in 1999 and 2002 than in 2007 and 2008 and were close to twice the sizes of single meals only in the latter period, probably because in the earlier years, when they were fed more frequently, chicks were unable to accept all the food provided by the second parent to return overnight.

The fact that parents performed fewer 2 to 3 d trips in 2007 and 2008 suggests that prey may have been present in sufficient quantity only at longer distances from the colony (Pichegru et al. 2007). This is consistent with a suspected northerly shift in the foraging areas of these birds in recent years (Guilford et al. 2008). Manx shearwaters prey mainly on 1 and 2 yr old clupeids (Brooke 1990), and so the negative effect of both increases in winter SST and decreases in the body masses of 2 yr old herring suggests that warming of surface waters may have had a negative impact on the growth and nutritional quality of prey, as also suggested for sandeels Amodytes marinus in the North Sea since the early 1980s (Wanless et al. 2004, 2005). In waters around western Britain, there is recent evidence that some fish species have also been affected through bottom-up processes (Todd et al. 2008). In our study, peak masses of chicks were negatively related to SST in the immediately preceding winter, suggesting that these effects have now been propagated to a higher trophic level, in a similar manner to that previously recorded in the North Sea (Frederiksen et al. 2004b) and in other oceanographic regions (Guinet et al. 1998, Gjerdrum et al. 2003).

In contrast to Manx shearwaters, guillemots Uria aalge and razorbills Alca torda (Alcidae) have apparently bred normally in recent years on Skomer (D. Boyle pers. comm.). This difference in responses may reflect differences in foraging depths, with guillemots and razorbills both being capable of diving much deeper than Manx shearwaters (Thaxter et al. 2010), or in the timing of breeding, since chicks of both these species have fledged from the breeding cliffs when Manx shearwaters are still feeding young chicks. However, in contrast to events during the breeding season, there is recent evidence that the over-winter survival of guillemots from Skomer is negatively related to the NAO index, and thus lower in warmer, wetter and windier winters, which have increased in frequency in recent years (Votier et al. 2005). Together, these 2 studies suggest that within the NE Atlantic region, adverse impacts of anthropogenic climate change on higher marine predators are no longer restricted to the vicinity of the North Sea.
LITERATURE CITED

- Ainley DG, Blight LK (2009) Ecological repercussions of historical fish extraction from the southern ocean. Fish Fish 10:13–38
- Hunt GL Jr, Stabeno P, Walters G, Sinclair E, Brodeur RD,


Editorial responsibility: Hans Heinrich Janssen,
Oldendorf/Luhe, Germany

Proofs received from author(s): January 25, 2011