Investigating the potential impacts of climate change on a marine turtle population

L. A. HAWKES*†, A. C. BRODERICK*, M. H. GODFREY‡ and B. J. GODLEY*

*Marine Turtle Research Group, Centre for Ecology and Conservation, School of Biosciences, University of Exeter, Cornwall Campus, Penryn TR10 9EZ, UK, †Bald Head Island Conservancy, PO Box 3109, 7000 Federal Road, Bald Head Island, NC 28461-7000, USA, ‡North Carolina Wildlife Resources Commission, 1507 Ann Street, Beaufort, NC 28516, USA

Abstract

Recent increases in global temperatures have affected the phenology and survival of many species of plants and animals. We investigated a case study of the effects of potential climate change on a thermally sensitive species, the loggerhead sea turtle, at a breeding location at the northerly extent of the range of regular nesting in the United States. In addition to the physical limits imposed by temperature on this ectothermic species, sea turtle primary sex ratio is determined by the temperature experienced by eggs during the middle third of incubation. We recorded sand temperatures and used historical air temperatures (ATs) at Bald Head Island, NC, to examine past and predict future sex ratios under scenarios of warming. There were no significant temporal trends in primary sex ratio evident in recent years and estimated mean annual sex ratio was 58% female. Similarly, there were no temporal trends in phenology but earlier nesting and longer nesting seasons were correlated with warmer sea surface temperature. We modelled the effects of incremental increases in mean AT of up to 7.5 $^{\circ}$ C, the maximum predicted increase under modelled scenarios, which would lead to 100% female hatchling production and lethally high incubation temperatures, causing reduction in hatchling production. Populations of turtles in more southern parts of the United States are currently highly female biased and are likely to become ultra-biased with as little as 1 °C of warming and experience extreme levels of mortality if warming exceeds 3 °C. The lack of a demonstrable increase in AT in North Carolina in recent decades coupled with primary sex ratios that are not highly biased means that the male offspring from North Carolina could play an increasingly important role in the future viability of the loggerhead turtle in the Western Atlantic.

Keywords: Caretta caretta, climate change, loggerhead turtle, temperature dependant sex determination, warming

Received 29 June 2006; revised version received 25 September 2006 and accepted 11 October 2006

Introduction

Global temperatures have warmed by approximately 0.6 °C in the last century at a rate that has been unprecedented in the last 1000 years [mean air and sea surface; Intergovernmental Panel on Climate Change (IPCC), 2001; Levitus *et al.*, 2001; Brohan *et al.*, 2006]. Modelled projections for climate change in the next 100 years predict further accelerated levels of warming between 0.3 and 7.5 °C for North America (IPCC, 2001; Zwiers, 2002; low and high emissions scenarios).

Correspondence: Dr Brendan J. Godley, tel. +44 (1326) 371861, fax +44 (1326) 371859, e-mail: bgodley@seaturtle.org

© 2007 The Authors Journal compilation © 2007 Blackwell Publishing Ltd However, this future climate change is expected to be heterogeneous, and not all areas will experience warming or similar rates of change in temperature. Therefore, biodiversity conservation strategies must increasingly look to the future and plan for potential outcomes over a range of possible changes in global, regional and local climate.

There is now a sizeable body of literature documenting advances in the annual phenology (timing of seasonal activities) of many animal and plant species in concert with observed climate change (see review in Walther *et al.*, 2002). Most commonly, spring events occur earlier (e.g. arrival at breeding grounds), reproductive seasons end earlier (Pike *et al.*, 2006) but there have also been records of later onset of autumnal events (e.g. delayed migration). Eighty-seven percent of species (from plants to vertebrates) reviewed by Parmesan & Yohe (2003) that exhibited shifts in phenology did so highly significantly in accord with climate change. For example, first plant flowering, tree budburst, arrival of migrant butterflies and birds, and timing of bird nesting have all occurred earlier with increasing spring air temperatures (ATs) (see review in Parmesan & Yohe, 2003). Arrival and departure dates of bird species in breeding grounds are often well recorded due to their visible nature and public appeal, and such dates have advanced for many short and medium distance migrants (Butler, 2003; Lehikoinen et al., 2004; Mills, 2005). Additionally, larger clutch sizes and faster offspring development have been documented in some European and North American bird species (Järvinen, 1989; McCleery & Perrins, 1998; Visser et al., 1998; Bergman, 1999; Brown et al., 1999; Winkler et al., 2002). In the marine environment, variations in temperatures directly drive abundance and distribution of many organisms. Seasonal peaks in copepod and zooplankton abundance have occurred up to 11 weeks earlier in response to increasing sea surface temperatures (SSTs; Greve et al., 2001), while migratory fish species have been moving to spring spawning grounds earlier (Carscadden et al., 1997; Sims et al., 2004).

In temperate zones, where cold winter temperatures often exceed physiological tolerance thresholds, many animals adapt by moving seasonally to more tolerable habitats to avoid stress or death. However, if temperatures were to increase with climate change, isotherms could shift poleward (Hughes, 2000), moving tolerable habitats, and therefore species distributions, towards higher latitudes (Shine *et al.*, 2002). Such range expansions have been recorded for many plant and animal species (see reviews in Walther *et al.*, 2002; Hickling *et al.*, 2006), with butterfly species, for example, exhibiting particularly significant range shifts to the order of several hundred kilometres north (Parmesan *et al.*, 1999; Davies *et al.*, 2006).

In addition to effects on phenology and distribution, climate change may have more direct impacts on thermally sensitive organisms (Walther *et al.*, 2002). In particular, ectotherms are sensitive to thermally available conditions that directly affect their performance, survival and reproduction. Many reptile species exhibit temperature-dependant sexual determination (TSD) where primary sex ratio is influenced by the temperature experienced by eggs during incubation. Even slight changes (\sim 1 °C) to incubation temperatures during this period may radically alter offspring sex ratios of TSD species (Janzen, 1994). For populations of species with TSD that have already been shown to produce highly

skewed primary sex ratios, climate change could further skew production of only one sex, for example, if no males are produced, females could lay unfertilized clutches, annual cohorts of hatchlings would be lost and could lead towards local extinctions.

In marine turtles, sex is determined by temperature in the middle third of incubation with female offspring produced at higher temperatures and males at lower temperatures within a thermal tolerance range of 25-35 °C (Ackerman, 1997). A mixture of sexes is produced within the threshold range of temperatures (TRT, approximately 4°C wide for the loggerhead sea turtle, Caretta caretta; Yntema & Mrosovsky, 1979) and 50% of either sex at the 'pivotal temperature' (around 29.0 °C, Mrosovsky, 1988). The loggerhead sea turtle is a species with TSD, mostly nesting on oceanic beaches outside the tropics, producing predominantly female hatchlings at most sites studied (Mrosovsky & Provancha, 1989, 1992; Mrosovsky, 1994; Marcovaldi et al., 1997; Hanson et al., 1998; Godley et al., 2001a, b; Oz et al., 2004) and is the most studied marine turtle species for TSD. In North America, loggerhead turtles are found nesting from southern Florida to southern Virginia. At the northern limit of the nesting range, where sand temperatures are cooler, a greater proportion of male offspring are thought to be produced in comparison with the highly female-biased sex ratios recorded in Florida (Mrosovsky et al., 1984; Mrosovsky & Provancha, 1992; Heppell *et al.*, 2003).

Under future scenarios of climate warming, and given the conservation concern regarding this species, it is important to widen our understanding of the potential effects of temperature increases on sea turtle populations and their potential to cope with such changes. Here, we review 26 years (1980–2005) of detailed data from a northerly nesting site for loggerhead turtles. In concert with six decades of local climate data we predict primary sex ratios, describe climate change patterns and investigate whether any shifts in phenology have occurred or are possible at this site. In addition, we estimate recent historical patterns in primary sex ratio and model possible future outcomes in the face of warming expected for turtles nesting at the northern portion, as well as further south, of the range.

Methods

Nesting data between May and October were recorded from 1980 to 2005 by a nightly sea turtle monitoring programme at Bald Head Island, NC, USA (33°50'N, 77°57'W, Fig. 1; see Hawkes *et al.*, 2005). Incubation duration was recorded as the number of days between egg laying and hatchling emergence at the sand surface. The duration of the nesting season (the number of days



Fig. 1 Map of Bald Head Island, NC, USA, showing the location of the Southport weather station and location of control sites for temperature data loggers (numbered circles 1–4).

between the first and last egg laying events) was calculated for each year as were the median and interquartile dates of nesting for each season.

Local AT data were obtained from the State Climate Office of North Carolina (http://www.nc-climate. ncsu.edu/services/request.html) for Southport, North Carolina (6.4 km northwest of study site, Fig. 1), from January 1940 to December 2005. Station instruments are maintained by the US National Weather Service and quality controlled by the US National Climatic Data Centre. Mean weekly SST data were sampled from the Met Office Hadley Centre's sea ice temperature and SST data sets (HadISST1) for 34.057°N and -77.189°W between 1980 and 2005. HadISST1 replaces the global sea ice and SST (GISST) data sets and contains *in situ* monthly sea surface observations and satellite derived estimates at the sea surface on a 1° latitude–longitude grid.

As pivotal temperature is a relatively conservative characteristic in sea turtles (Freedberg & Wade, 2001), typically varying <1.8 °C around 29.2 °C for the loggerhead sea turtle (Ackerman, 1997), we used published models for loggerhead turtles from this population

relating incubation temperature to proportion of female hatchlings produced (Mrosovsky, 1988) to estimate past and present hatchling sex ratios. As we were not able to record individual nest temperatures for the study period we modelled, nest temperatures as follows: sand temperatures were recorded during the 2003 nesting season using four calibrated temperature data loggers (Gemini data loggers, Chichester, UK) buried at 45 cm depth (mean nest depth for Bald Head Island, L. A. Hawkes, unpublished data) at four control locations on the beach (Fig. 1) and hourly temperatures were collected from 5 July until 14 November 2003. We examined the relationship between mean daily sand temperature and mean daily AT collected at Southport for 2003 (highly significantly related: sand temperature = 0.72 AT + 9.60, $R^2 = 0.80$; regression $F_{1,131} = 513.37$, P < 0.001) and used this to model sand temperatures for the study period (1980-2005). Metabolic heating of the clutch during incubation must be considered (Broderick et al., 2001) and therefore we added an estimate of metabolic heating of loggerhead turtle eggs from Florida, USA in the middle third period of incubation (Hanson et al., 1998; 1.1 °C) to predict nest

temperatures in the middle third of incubation for the study period (1980–2005).

Further, in order to heuristically model the possible effects of climate change on primary sex ratios, we added the resulting increases in sand temperature from predicted increases in AT of 1–7.5 °C (the predicted range of maximum rise in global temperature by 2100; Houghton, 2001) to the estimated nest temperatures 1980–2005 and modelled the resulting primary sex ratios. We repeated the same process for half-month average sand temperature recorded for a Florida nesting beach (1986–1990, Cape Canaveral Air Force Station; Mrosovsky & Provancha, 1992), using the deepest sand temperature data recorded (60 cm) on the low shore, to parsimoniously generate the minimum plausible temperatures and least female biased estimates.

Results

Long-term mean ATs at Southport, NC (1940–2005), have decreased significantly over the 64 years (by approximately 1.7 °C, regression: $F_{1,62} = 13.88$, P < 0.01, $R^2 = 0.18$, Fig. 2a). The same significant decrease is evident if temperatures during the nesting season of each year are sampled (approximately 1.9 °C decrease, 4 May to 15 November 1940–2005, regression: $F_{1,63} = 14.79$, P < 0.01, $R^2 = 0.19$, Fig. 2b). During the study period (1980–2005) annual SST was significantly related to annual AT (Pearsons r = 0.452, n = 26, AT = 1.3 SST °C -13.5, P < 0.05). Although SST warmed significantly (regression: annual $F_{1,25} = 12.31$, P < 0.01, $R^2 = 0.34$, seasonal $F_{1,25} = 22.89$, P < 0.01, $R^2 = 0.49$, Fig.

2c and d), AT did not (regression: annual $F_{1,25} = 1.47$, P > 0.05, $R^2 = 0.06$, seasonal $F_{1,25} = 2.05$, P > 0.05, $R^2 = 0.08$) and we highlight this as a caveat of this type of study. There were also no trends in ATs in the month preceding the nesting season (April regression: $F_{1,24} = 0.56$, P > 0.05, $R^2 = 0.02$).

Given the predicted climate change by the IPCC (2001), we investigated metrics of phenology in relation to inter annual variability in environmental conditions to see whether changes in nesting patterns were driven by climate. Because sea turtles experience ambient temperature through the water rather than air, we investigated relationships with SSTs. In years with warmer spring (April and May) SSTs, nesting started significantly earlier (mean April SST vs. first nest regression: $F_{1,22} = 4.72$, P < 0.05, first nest = -3.9 SST °C + 223.5, $R^2 = 0.18$, 12 days earlier per 1 °C and mean May SST vs. first nest regression: $F_{1,22} = 8.20$, P < 0.01, first nest = -5.2 SST °C + 265.2, $R^2 = 0.28$, 18.3 days earlier per 1 °C, Fig. 3). This increased the duration of the nesting season significantly in warmer years (April regression $F_{1,22} = 7.15 P < 0.05$, duration = 4.8 SST °C-14.5, $R^2 = 0.25$, 18.3 days longer per 1 °C, May regression $F_{1,22} = 3.47$, *P* < 0.05, duration = 5.1 SST °C-34.8, $R^2 = 0.14$, 36 days longer per 1 °C) but the date of last nest laid was not significantly related to any metric of SST. Although we have demonstrated the link among environment and phenology and a significant increase in SSTs over the study period, there was no significant temporal trend in the date of first nest laid or duration of the nesting season (first nest regression $F_{1,22} = 2.26$, P > 0.05, $R^2 = 0.1$; duration regression $F_{1,22} = 0.36$,



Fig. 2 Long-term trends in temperature: Annual deviation from (a) long-term mean annual air temperature (17.28 °C) and (b) long-term mean seasonal air temperature (May–November each year; 22.70 °C) recorded at Southport, NC, USA, between 1940 and 2005 and (c) long-term mean annual sea surface temperature (SST; 23.23 °C) and (d) long-term mean seasonal SST (25.93 °C) recorded at Cape Fear, NC between 1980 and 2005. The study period (1980–2005) is indicated in parts (a) and (b) by the horizontal arrow.



Fig. 3 Relationship between first nest laid each year and mean a) April sea surface temperature (SST) and (b) May SST at Bald Head Island 1983–2005. Data for 1980–1982 have been excluded due to incomplete records in nesting metrics.

P > 0.05, $R^2 = 0.04$). None of the other phenological attributes (date of latest hatchling emergence, lower and upper quartiles and median egg-laying date) were significantly related to air or SST over the temporal period studied.

The estimated mean sex ratio for the study period (calculated from modelled nest temperatures) (Fig. 4) was 58% female hatchlings. There was no significant temporal change in annual sex ratio production estimated using modelled incubation temperature (1980-2005, regression: $F_{1,23} = 1.12$, P > 0.05, $R^2 = 0.05$). With the predicted rise in future AT, nests will begin to incubate at sand temperatures above the pivotal temperature (29.2 °C; Mrosovsky, 1988) for loggerheads, producing more female offspring. Simple heuristic simulation of likely hatchling sex ratios under these conditions (Fig. 5a and b) show that over 80% female hatchling production is predicted to occur in this part of North Carolina with 2 °C increase in AT. Further south, at Cape Canaveral, Florida, total feminization would occur with 2 °C of warming in AT and after 3 °C clutches would begin to incubate at temperatures above the lethal limit (35°C; Ackerman, 1997). However, such clutch death is not likely to begin until 5 °C of warming in AT has occurred in North Carolina. With the maximum 7.5 °C of predicted warming in AT, more than



Fig. 4 Frequency histogram of modelled mean temperatures experienced by nests at Bald Head Island (1980–2005) during the middle third of incubation, including an estimate of metabolic heating. Pivotal incubation temperature (29.2 °C) is shown as vertical dashed line and with male biased and female biased incubation temperatures indicated.



Fig. 5 Ratios of resultant sex of hatchlings at present and under varying predictions of future climate change for (a) Bald Head Island, NC and (b) Cape Canaveral, Florida, 1986–1990 showing proportion female (grey) and proportion male (white). Black signifies the expected mortality resulting from incubation over the upper tolerance limit ($35 \,^{\circ}$ C) in response to increases in air temperature between 1 and 7.5 °C, where the final bar shows 7.5 °C of warming.

40% of clutches at our study site and 100% of clutches at Cape Canaveral, will incubate at temperatures above the lethal limit, suggesting that nest incubation would



Fig. 6 Thermal niche available for incubation, showing current (solid line) and modelled (broken lines) sand temperatures (primary *y*-axis) resulting from increases of 2, 4 and 7.5 °C in air temperature (sand temperature = 0.7154 air temperature +9.6023). Black bars show proportion of clutches incubating in each week of the season (1980–2005, secondary *y*-axis). Shaded area represents the upper thermal limit (UTL) for incubation (33–35 °C; Ackerman, 1997).

no longer be viable at some Florida beaches during the current seasonal window of nesting. However, with climate warming, the actual thermal niche available to turtles for nesting will become wider (as a very simplistic example, with a mean increase of $7.5 \,^{\circ}$ C AT, the worst case scenario of warming, the suitable thermal conditions for incubation almost doubles, Fig. 6), providing sea turtles the opportunity to alter temporal nesting distribution to earlier and/or later months to mitigate increased incubation temperatures.

Discussion

The climate system of North Carolina is one of the most complex in North America (Boyles & Raman, 2003). The proximity of the Gulf Stream to coastal North Carolina acts as an ameliorating influence to climatic variations and combines with local topography to produce a highly variable regional climate that can differ substantially to climate patterns observed elsewhere on the south eastern coast of the United States, although 10 of the last 12 years have been above the long-term average. North Carolina has not yet experienced pronounced warming and it may be that the study area is less likely to experience the significant warming expected elsewhere in the Northern Hemisphere by the IPCC. Boyles & Raman (2003) found that average ATs were warmest during a period between 1949 and 1958 and decreased slightly after. This is consistent with the gradual cooling trend we observed in AT at Southport since 1940.

Our results, based on 26 years of data, suggest that loggerhead nests in North Carolina currently produce

proportionally more males (42% males) than nests laid in Florida (typically <10% males; Mrosovsky & Provancha, 1989). These results are in accord with previous studies of sex ratios from the northern nesting sites in the United States (Mrosovsky *et al.*, 1984; Webster & Gouveia, 1989). This is one of only a few studied loggerhead rookeries that are not predominantly female producing, others located in Espirito Santo, Brazil (Marcovaldi *et al.*, 1997; Baptistotte *et al.*, 1999) and Heron Island, Australia (Limpus *et al.*, 1983). Taken in concert with the environmental data sampled in this study, it appears that the primary sex ratio in North Carolina is not likely to become radically female biased in future years unless warming is severe.

However, given that Florida could likely experience warming (IPCC, 2001) and is not buffered by oceanic features as in North Carolina (Boyles & Raman, 2003), there is a real possibility of further skewing or even complete feminization of the primary sex ratios in Florida, which holds the vast majority of nesting in the United States (Shoop & Kenney, 1992). The results of our simple modelling show that 2 °C increase in ATs (within the minimum expected warming scenario of the IPCC, 2001) is sufficient to result in total feminization and with 3 °C of warming, many nests in southern Florida would begin to experience incubation temperatures above lethal limits. Therefore, in addition to extreme primary sex ratios, decreased or zero hatch success is worthy of consideration for all nesting populations with current highly skewed sex ratios (e.g. Mediterranean: Broderick et al., 2000; Godley et al., 2001a, b; Öz et al., 2004; Florida: Mrosovsky & Provancha, 1992; Mrosovsky, 1994; Hanson et al., 1998; Brazil: Marcovaldi et al., 1997). However, it should be noted that primary sex ratios may not reflect those of the adult population, although in-water studies suggest that female biases are maintained in juvenile and sub-adult loggerhead turtles (Wibbels et al., 1987, 1991; Owens, 1997).

If warming occurs, turtles from northern and southern beaches could prevent extremely skewed sex ratios by (a) adapting *in situ* by adjusting their pivotal temperature, (b) altering nesting spatially (e.g. latitude, height on the beach, depth or substrate type, Hays *et al.*, 2001) or (c) altering nesting temporally (e.g. nesting in cooler periods at the beginning and end of the season in order to exploit incubation environments likely to produce more males; Mrosovsky *et al.*, 1984; Naro-Maciel *et al.*, 1999; Hays *et al.*, 2003; Pike *et al.*, 2006). Pivotal temperature is a relatively conservative TSD character in sea turtles (Mrosovsky, 1988; Freedberg & Wade, 2001; Godfrey & Mrosovsky, 2006) with little variation observed across the wide-ranging geographic locations and latitudes over which it has been studied (e.g. Wibbels, 2003; Godfrey & Mrosovsky, 2006; but see studies in fresh water turtles e.g. Ewert *et al.*, 2005). We, therefore, consider it unlikely that turtles would be able to adapt pivotal temperature in the shortterm (less than several hundred years; cf. Morjan, 2003; Glen & Mrosovsky, 2004; Nelson *et al.*, 2004), although plasticity in the transitional range of temperatures surrounding the pivotal temperature could potentially be adaptive (Girondot, 1999).

The ability to adapt spatially in response to thermally variable habitat occurs in some reptile species where females have been shown to consistently select nest microenvironments with similar thermal conditions, even in different locations (e.g. the painted turtle, Chrysemys picta, Janzen & Morjan, 2001; and the hawksbill turtle, Eretmochelys imbricata, Kamel & Mrosovsky, 2005). For example, at warmer sites, female painted turtles seek more shaded nest locations, and can thus manipulate the thermal habitat, and potentially the sex ratio, of their offspring (but see Morjan, 2003). However, as loggerhead turtles usually maintain a degree of nest site fidelity, returning to nest within 0-5 km of previous nests both within and between nesting seasons (Miller et al., 2003) and climate warming would be expected to occur over a large (regional) scale, the different scales over which change could occur could prevent adaptation by turtles shifting nesting location en masse. Loggerhead turtles could, however, alter the specific nest environment by nesting, as Hawksbill turtles do, in cooler microclimates, such as in areas shaded by vegetation (Kamel & Mrosovsky, 2006) or closer to the sea, (Foley et al., 2006), and natal homing of newly recruiting nesting turtles could contain a margin of geographical error that could mitigate philopatry that cannot be ruled out at this point. In addition, as more females are produced in the larger southern subpopulation, the frequency of nesting should be expected to increase in the south relative to the north due to philopatry (Morjan, 2003).

Loggerhead turtles at Bald Head Island currently nest throughout the available temporal window that provides a tolerable environment for nests (Fig. 6) but the actual niche available for turtle nesting under future climate scenarios could be very different. Phenological correlations in this study and others (Mrosovsky *et al.*, 1984; Weishampel *et al.*, 2004; Pike *et al.*, 2006) suggest that either changing temperatures in the future could drive adaptive phenological change to maintain current thermal conditions in nests or that turtles may adapt the temporal spread of nesting to suit the available thermal niche. However, it should be noted that Pike *et al.* (2006) demonstrated that earlier seasons can also end earlier, probably because sea turtles are capital breeders (Bonnet *et al.*, 1998), meaning that the fundamental and realised thermal niches for nesting may not necessarily be the same. Because the distribution of nesting has a low variance (Fig. 6), our model shows that turtles would have to alter the temporal distribution of nesting very little (by only a few days) to adapt to $1 \,^{\circ}$ C of warming, and by up to a week for $3 \,^{\circ}$ C of warming (worst expected under lower case emissions scenario; IPCC, 2001). This strategy could, therefore, offer the most likely adaptive mechanism available to marine turtles in response to climate change.

However, relying on accumulated energy reserves for breeding, they typically make long migratory journeys to get from foraging to breeding areas. Turtles, therefore, must not only have stored energy from the foraging grounds (which may be some 800 km away, L. A. Hawkes et al., in press) for reproduction before the nesting season, but environmental conditions at the foraging grounds likely influence vitellogenesis and may therefore constrain the timing of migration and breeding, thus complicating phenological adaptation. Although we have described earlier nesting in years with warmer SST, this may reflect more rapid egg production following mating as oogenesis is likely driven by prevailing environmental temperature (Hays et al., 2002; Hamann et al., 2003), and females are thought to migrate to the nesting beach ca. 1 month before beginning laying (Godley et al., 2002). Because capital breeding constrains reproductive output (loggerhead turtles normally lay two to four clutches per nesting season, maximum seven clutches; Schroeder et al., 2003), even if turtles started to nest earlier in the season, later clutches would still incubate in the period of very high temperatures (above thermal tolerance with maximum levels of warming) and turtles are unlikely to still be nesting in the later, cooler part of the season (Pike et al., 2006).

Irrespective, the fact that many loggerhead sea turtle populations that already exhibit heavily female-biased primary sex ratios may be negatively impacted by climate warming merits further investigation. The potentially disproportionate importance of male producing beaches (e.g. Mrosovsky et al., 1984; Baptistotte et al., 1999) that can interbreed with female producing rookeries (Bowen et al., 2005) needs to be carefully examined, particularly nesting beaches such as Bald Head Island that not only exhibit more balanced sex ratios but appear less likely to be affected by climate change. Emphasis should be placed on maintaining suitable nesting habitat for sea turtles in North Carolina and other regions that are important for male production. Anthropogenic activities likely to alter the thermal properties of naturally available nesting habitat should be given careful consideration. In particular, beach

'nourishment' programs (which are becoming increasingly widespread) that deposit sand on eroded beaches along the oceanic coastline of the United States to protect beachfront property (Crain et al., 1995; Milton et al., 1997; Rumbold et al., 2001; Peterson & Bishop, 2005) often use sand that differs radically from natural sands and are often darker in colour. Nests in darker sand incubate at higher temperatures producing more females (Hays et al., 2001). Moreover, although turtles have evolved with continuously eroding and accreting oceanic beaches, the combination of renourishment programs, along with other anthropogenic threats such as beachfront development, and predicted sea level rise (from 0.11 to 0.43 mm yr^{-1} ; IPCC, 2001) could cause a 'coastal squeeze', where landward movement of beaches is prevented by physical (anthropogenic) barriers such as beachfront development (Fish et al., 2005). In addition, increased storm surges and coastal land loss will threaten beaches backed by salt marshes, such as those making up the majority of the outer banks of North Carolina.

The sex ratios generated in this model should be tempered with elements of caution; actual primary sex ratios from sea turtle nesting beaches can be influenced by numerous factors including (but not limited to) humidity, rainfall, tidal inundation, the total number of eggs incubating, egg position in nest (Carthy *et al.*, 2003), nest relocation, nest substrate, hatching success, egg loss and intra-seasonal variation in temperatures and were not controlled for in the model. Rather, the results of this study shed light on potential outcomes if scenarios envisaged under climate warming do occur. Note that climate change (and sea level rise) is not expected to be geographically uniform (IPCC, 2001) and it is likely that some areas may not experience the profound warming predicted.

Even though climate change models are subject to controversy and the sex ratios generated in this study are based on a simplistic model, there are some important take home messages from this study. That there are nesting beaches that are both male producing and somewhat shielded from future climate change is noteworthy and the thermal conditions of these beaches should be protected from anthropogenic activities likely to alter the natural state of the incubation environment. The highly female-skewed southern Florida population may, under worst case scenarios, reach a 100% female asymptote, at which point males from further north in the population may be essential for the continued production of hatchlings from southern beaches. Finally, with a few degrees of warming, large proportions of the clutches of the largest loggerhead nesting population in the world may begin to experience high levels of thermal mortality.

Acknowledgements

Many thanks to the staff of North Carolina Wildlife Resources Commission for storing and managing data, Melissa Hedges, staff, students and supporters of the Bald Head Island Conservancy, Michael Coyne and Matthew Witt for help with data products and Fiona Glen for help with temperature loggers. The State Climate Office of North Carolina provided air temperature data. L. A. H. is supported by the Anning Morgan Bursary awarded to the Centre for Ecology and Conservation at the University of Exeter in Cornwall. B. J. G. and A. C. B. are supported by European Social Fund Project number 041015SW1, Natural Environment Research Council Fellowship, Foreign and Commonwealth Environment Fund for the Overseas Territories, Darwin Initiative and the Overseas Territories Environment Programme. This manuscript benefited greatly from constructive comments from four anonymous reviewers.

References

- Ackerman RA (1997) The nest environment and embryonic development of sea turtles. In: *The Biology of Sea Turtles* (eds Lutz PL, Musick JA), pp. 83–106. CRC Press, Boca Raton.
- Baptistotte C, Scalfoni JT, Mrosovsky N (1999) Male-producing thermal ecology of a southern loggerhead turtle nesting beach in Brazil: implications for conservation. *Animal Conservation*, 2, 9–13.
- Bergman E (1999) Long term increases in numbers of earlyfledged reed warblers (*Acrocephalus scirpaceus*) at lake Constance (Southern Germany). *Journal of Ornithology*, **140**, 80–86.
- Bonnet X, Bradshaw D, Shine R (1998) Capital versus income breeding: an ectothermic perspective. Oikos, 83, 333–342.
- Bowen BW, Bass AL, Soares L, Toonen RJ (2005) Conservation implications of complex population structure: lessons from the loggerhead turtle. *Molecular Ecology*, 14, 2389–2402.
- Boyles RP, Raman S (2003) Analysis of climate trends in North Carolina (1949–1998). Environment International, 29, 263–275.
- Broderick AC, Godley BJ, Hays GC (2001) Metabolic heating and the prediction of sex ratios for green turtles. *Physiological and Biochemical Zoology*, 74, 161–170.
- Broderick AC, Godley BJ, Reece S, Downie JR (2000) Incubation periods and sex ratios of green turtles: highly female biased hatchling production in the eastern Mediterranean. *Marine Ecology Progress Series*, **202**, 273–281.
- Brohan PB, Kennedy JJ, Harris I, Tett SFB, Jones PD (2006) Uncertainty estimates in regional and global observed temperature changes: a new dataset from 1850. *Journal of Geophy*sical Research, **111**, D12106.
- Brown JL, Li SH, Bhagabati N (1999) Long term trend toward earlier breeding in an American bird: a response to global warming? *Proceedings of the National Academy of Sciences of the* USA, **96**, 5565–5569.
- Butler CJ (2003) The disproportionate effect of global warming on the arrival dates of short-distance migratory birds in North America. *Ibis*, **145**, 484–495.
- Carscadden J, Nakashima BS, Frank KT (1997) Effects of fish length and temperature on the timing of peak spawning in capelin (*Mallotus villosus*). *Canadian Journal of Fisheries and Aquatic Sciences*, **54**, 781–787.

© 2007 The Authors

- Carthy RR, Foley AM, Matsuzawa Y (2003) Incubation environment of loggerhead turtle nests. In: *Loggerhead Sea Turtles* (eds Bolten AB, Witherington BE), pp. 144–153. Smithsonian Books, Washington.
- Crain DA, Bolten AB, Bjorndal KA (1995) Effects of beach renourishment on sea turtles: review and research initiatives. *Restoration Ecology*, **3**, 95–104.
- Davies ZG, Wilson RJ, Coles S, Thomas CD (2006) Changing habitat associations of a thermally constrained species, the silver-spotted skipper butterfly, in response to climate warming. *Journal of Animal Ecology*, **75**, 247–256.
- Ewert MA, Lang JW, Nelson CE (2005) Geographic variation in the pattern of temperature-dependant sex determination in the American snapping turtle (*Chelydra serpentina*). *Journal of the Zoological Society of London*, **265**, 81–95.
- Fish MR, Côté IM, Gill JA, Jones AP, Renshoff S, Watkinson AR (2005) Predicting the impact of sea-level rise on Caribbean sea turtle nesting habitat. *Conservation Biology*, **19**, 482–491.
- Foley AM, Peck SA, Harman GR (2006) Effects of sand characteristics and inundation on the hatching success of loggerhead sea turtle (*Caretta caretta*) clutches on low-relief mangrove islands in southwest Florida. *Chelonian Conservation and Biology*, **5**, 32–41.
- Freedberg S, Wade MJ (2001) Cultural inheritance as a mechanism for population sex-ratio bias in reptiles. *Evolution*, **55**, 1049–1055.
- Girondot M (1999) Statistical description of temperature-dependent sex determination using maximum likelihood. *Evolutionary Ecology Research*, **1**, 479–486.
- Glen F, Mrosovsky N (2004) Antigua revisited: the impact of climate change on sand and nest temperatures at a hawksbill turtle (*Eretmochelys imbricata*) nesting beach. *Global Change Biology*, **10**, 2036–2045.
- Godfrey MH, Mrosovsky N (2006) Pivotal temperature for green turtles, *Chelonia mydas*, nesting in Suriname. *Herpetological Journal*, 16, 55–61.
- Godley BJ, Broderick AC, Downie JR *et al.* (2001b) Thermal conditions in nests of loggerhead turtles: further evidence suggesting female skewed sex ratios of hatchling production in the Mediterranean. *Journal of Experimental Marine Biology and Ecology*, **263**, 45–63.
- Godley BJ, Broderick AC, Frauenstein R, Glen F, Hays GC (2002) Reproductive seasonality and sexual dimorphism in green turtles. *Marine Ecology Progress Series*, **226**, 125–133.
- Godley BJ, Broderick AC, Mrosovsky N (2001a) Estimating hatchling sex ratios of loggerhead turtles in Cyprus from incubation durations. *Marine Ecology Progress Series*, **210**, 195– 201.
- Greve W, Lange U, Reiners F, Nast J (2001) Predicting the seasonality of North Sea zooplankton. *Senckenbergiana Maritime*, **31**, 263–268.
- Hamann M, Limpus CJ, Owens DW (2003) Reproductive cycles of males and females. In: *Biology of Sea Turtles*, Vol. 2 (eds Lutz PL, Musick JA), pp. 275–306. CRC Press, Boca Raton.
- Hanson J, Wibbels T, Martin RE (1998) Predicted female bias in sex ratios of hatchling loggerhead sea turtles from a Florida nesting beach. *Canadian Journal of Zoology*, **76**, 1850–1861.

- Hawkes LA, Broderick AC, Godfrey MH, Godley BJ (2005) Status of nesting loggerhead turtles at Bald Head Island (North Carolina, USA) after 24 years of intensive monitoring and conservation. *Oryx*, **39**, 65–72.
- Hays GC, Ashworth JS, Barnsley MJ *et al.* (2001) The importance of sand albedo for the thermal conditions on sea turtle nesting beaches. *Oikos*, **93**, 87–94.
- Hays GC, Broderick AC, Glen F, Godley BJ, Houghton JDR, Metcalfe JD (2002) Water temperature and internesting intervals for loggerhead (*Caretta caretta*) and green (*Chelonia mydas*) sea turtles. *Journal of Thermal Biology*, **27**, 429–432.
- Hays GC, Broderick AC, Glen F, Godley BJ (2003) Climate change and sea turtles: a 150 year reconstruction of incubation temperatures at a major marine turtle rookery. *Global Change Biology*, 9, 642–646.
- Heppell SS, Snover ML, Crowder LB (2003) Sea turtle population ecology. In: *Biology of Sea Turtles*, Vol. 2 (eds Lutz PL, Musick JA), pp. 275–306. CRC Press, Boca Raton.
- Hickling R, Roy DB, Hills JK, Fox R, Thomas CD (2006) The distributions of a wide range of taxonomic groups are expanding polewards. *Global Change Biology*, **12**, 450–455.
- Houghton JR (2001) *Climate Change 2001: The Science of Climate Change.* Cambridge University Press, New York.
- Hughes L (2000) Biological consequences of global warming: is the signal already apparent? *Trends in Ecology and Evolution*, **15**, 56–61.
- Intergovernmental Panel on Climate Change (IPCC) (2001) Climate Change 2001: The Scientific Basis. Contribution of working group I to the third assessment report of the IPCC (eds Houghton JT, Ding Y, Griggs DJ, Noguer M, van der Linden PJ, Xiaosu D), Cambridge University Press, Cambridge, 881 pp.
- Janzen FJ (1994) Climate change and temperature-dependent sex determination in reptiles. *Proceedings of the National Academy of Sciences of the USA*, **91**, 7487–7490.
- Janzen FJ, Morjan CL (2001) Repeatability of microenvironmentspecific nesting behaviour in a turtle with environmental sex determination. *Animal Behaviour*, **62**, 73–82.
- Järvinen A (1989) Patterns and causes of long-term variation in reproductive traits of the pied fly-catcher *Ficedula hypoleuca* in Finnish Lapland. *Ornis Fennica*, **66**, 24–31.
- Kamel SJ, Mrosovsky N (2005) Repeatability of nesting preferences in the hawksbill sea turtle, *Eretmochelys imbricata*, and their fitness consequences. *Animal Behaviour*, **70**, 819–828.
- Kamel SJ, Mrosovsky N (2006) Deforestation: risk of sex ratio distortion in hawksbill sea turtles. *Ecological Applications*, 16, 923–931.
- Lehikoinen E, Sparks TH, Zalakevicius M (2004) Arrival and departure dates. *Birds Climate Change*, **35**, 1–31.
- Levitus S, Antonov JI, Wang J, Delworth TL, Dixon KW, Broccoli AJ (2001) Anthropogenic warming of earth's climate system. *Science*, **292**, 267–270.
- Limpus CJ, Reed P, Miller JD (1983) The influence of choice of nesting beach on sex ratio. In: *Proceedings Inaugural Great Barrier Reef Conference* (eds Baker JJ, Carter RM, Sammarco PW, Stark KP), pp. 397–402. James Cook University Press, North Queensland.

© 2007 The Authors

Journal compilation © 2007 Blackwell Publishing Ltd, Global Change Biology, doi: 10.1111/j.1365-2486.2006.01320.x

- Marcovaldi MA, Godfrey MH, Mrosovsky N (1997) Estimating sex ratios of loggerhead turtles in Brazil from pivotal incubation durations. *Canadian Journal of Zoology*, **75**, 755–770.
- McCleery RH, Perrins CM (1998) Temperature and egg laying trends. *Nature*, **391**, 30–31.
- Miller JD, Limpus CJ, Godfrey MH (2003) Nest site selection, oviposition, eggs, development, hatching, and emergence of loggerhead turtles. In: *Loggerhead Sea Turtles* (eds Bolten AB, Witherington BE), pp. 125–143. Smithsonian Books, Washington.
- Mills AM (2005) Changes in the timing of spring and autumn migration in North American migrant passerines during a period of global warming. *Ibis*, **147**, 259–269.
- Milton SL, Schulman AA, Lutz PL (1997) The effect of beach nourishment with aragonite versus silicate sand on beach temperature and loggerhead sea turtle nesting success. *Journal* of *Coastal Research*, **13**, 904–915.
- Morjan CL (2003) How rapidly can maternal behavior affecting primary sex ratio evolve in a reptile with environmental sex determination? *American Naturalist*, **162**, 205–219.
- Mrosovsky N (1988) Pivotal temperatures for loggerhead turtles (*Caretta caretta*) from northern and southern nesting beaches. *Canadian Journal of Zoology*, **66**, 661–669.
- Mrosovsky N (1994) Sex ratios of sea turtles. *The Journal of Experimental Zoology*, **270**, 16–27.
- Mrosovsky N, Hopkins-Murphy SR, Richardson JI (1984) Sex ratios of sea turtles: seasonal changes. *Science*, **225**, 739–741.
- Mrosovsky N, Provancha J (1989) Sex ratio of loggerhead sea turtles hatching on a Florida beach. *Canadian Journal of Zoology*, 67, 2533–2539.
- Mrosovsky N, Provancha J (1992) Sex ratio of hatchling loggerhead sea turtles: data and estimates from a 5-year study. *Canadian Journal of Zoology*, **70**, 530–538.
- Naro-Maciel E, Mrosovsky N, Marcovaldi MA (1999) Thermal profiles of sea turtle hatcheries and nesting areas at Praia do Forte, Brazil. *Chelonian Conservation and Biology*, **3**, 407–413.
- Nelson MJ, Thompson MB, Pledger S, Keall SN, Daugherty CH (2004) Do TSD, sex ratios, and nest characteristics influence the vulnerability of tuatara to global warming? *International Con*gress Series, **1275**, 250–257.
- Owens DW (1997) Hormones in the life history of sea turtles. In: *The Biology of Sea Turtles* (eds Lutz PL, Musick JA), pp. 315–341. CRC Press, Boca Raton.
- Öz M, Erdoğan A, Kaska Y et al. (2004) Nest temperatures and sex-ratio estimates of loggerhead turtles at Patara beach on the south-western coast of Turkey. *Canadian Journal of Zoology*, 82, 94–101.
- Parmesan C, Ryrholm N, Stefanescu C *et al.* (1999) Poleward shifts in geographical ranges of butterfly species associated with regional warming. *Nature*, **399**, 579–583.
- Parmesan C, Yohe G (2003) A globally coherent fingerprint of climate change impacts across natural systems. *Nature*, 421, 37–42.

- Peterson CH, Bishop MJ (2005) Assessing the environmental impacts of beach nourishment. *BioScience*, **55**, 887–896.
- Pike DA, Antworth RL, Stiner JC (2006) Earlier nesting contributes to shorter nesting seasons for the Loggerhead sea turtle, *Caretta caretta. Journal of Herpetology*, **40**, 91–94.
- Rumbold DG, Davis PW, Perretta C (2001) Estimating the effect of beach nourishment on *Caretta caretta* (Loggerhead sea turtle) nesting. *Restoration Ecology*, **9**, 304–310.
- Schroeder BA, Foley AM, Bagley DA (2003) Nesting patterns, reproductive migrations, and adult foraging areas of loggerhead turtles. In: *Loggerhead Sea Turtles* (eds Bolten AB, Witherington BE), pp. 114–124. Smithsonian Books, Washington.
- Shine R, Barrott EG, Elphick MJ (2002) Some like it hot: effects of forest clearing on nest temperatures of montane reptiles. *Ecology*, 83, 2808–2815.
- Shoop CR, Kenney RD (1992) Seasonal distributions and abundances of loggerhead and leatherback sea turtles in waters of the northeastern United States. *Herpetological Monographs*, **6**, 43–67.
- Sims DW, Wearmouth VJ, Genner MJ, Southward AJ, Hawkins SJ (2004) Low-temperature driven early spawning of a temperate marine fish. *Journal of Animal Ecology*, **73**, 333–341.
- Visser ME, van Noordwijk AJ, Tinbergen JM, Lessells CM (1998) Warmer springs lead to mistimed reproduction in great tits (*Parus major*). *Proceedings of the Royal Society of London B*, **265**, 1867–1870.
- Walther GR, Post E, Convey P *et al.* (2002) Ecological responses to recent climate change. *Nature*, **416**, 389–395.
- Webster WD, Gouveia JF (1989) Predicting hatchling sex ratios in loggerhead sea turtles (*Caretta caretta*) by incubation duration. In: *Proceedings of the Eighth Annual Workshop on Sea Turtle Conservation and Biology* (compiler Schroeder B), pp. 127–128. NOAA Technical Memorandum NMFS-SEFSC 214.
- Weishampel JF, Bagley DA, Ehrhart LM (2004) Earlier nesting by loggerhead sea turtles following sea surface warming. *Global Change Biology*, **10**, 1–4.
- Wibbels T (2003) Critical approaches to sex determination in sea turtles. In: *Biology of Sea Turtles*, Vol. 2 (eds Lutz PL, Musick JA, Wyneken J), pp. 103–134. CRC Press, Boca Raton.
- Wibbels T, Martin RE, Owens DW, Amoss MS (1991) Femalebiased sex ratios of immature loggerhead sea turtles inhabiting the Atlantic coastal waters of Florida. *Canadian Journal of Zoology*, 69, 2973–2977.
- Wibbels T, Owens DW, Morris YA, Amoss MS (1987) Sexing techniques and sex ratios for immature loggerheads captured along the Atlantic coast of the United States. In: *Ecology of East Florida Sea Turtles* (ed. Witzell WN), pp. 65–74. NOAA Technical Report, NMFS-53.
- Winkler DW, Dunn PO, McCulloch CE (2002) Predicting the effects of climate change on avian life-history traits. *Proceedings* of the National Academy of Science, **15**, 13595–13599.
- Yntema CN, Mrosovsky N (1979) Incubation temperature and sex ratio in hatchling loggerhead turtles: a preliminary report. *Marine Turtle Newsletter*, **11**, 9–10.
- Zwiers FW (2002) The 20 year forecast. Science, 416, 690-691.