INTRODUCTION

Sea turtles are long-lived, late maturing species that spend virtually all their lives in marine waters. All but one species are listed globally as endangered or critically endangered (IUCN 2006). Historically, most sea turtle research has focused on the brief emergences of nesting females. However, unless the populations are at equilibrium, counts of nesting turtles are an imperfect index of the current status of the entire population because such counts reflect nest production decades earlier. The effects of low production, an emerging source of mortality, or conversely, the removal of a source of mortality may not be detected on the beaches for decades. A few long-term studies of sea turtles on foraging grounds have provided valuable insight into population structure and trends. For example, work in Australia (Chaloupka & Limpus 2001) and the Bahamas (Bjorndal et al. 2005) provided information on trends in the abundance of green turtle Chelonia mydas populations in 2 ocean basins.

Sea turtle life histories are complex and comprise several life stages. Over their lifetime, sea turtles may inhabit virtually an entire ocean basin, including oceanic and coastal waters. Furthermore, sea turtle subpopulations each potentially have a different abundance trend and apparently mix non-randomly on the neritic foraging grounds (Broderick et al. 1994, Sears et al. 1995, Bolten et al. 1998, Bowen et al. 2004). Thus, multiple foraging grounds, with potentially differential demographics, contribute to a given nesting assemblage. Given such complexity, it is unlikely that estimating absolute abundance would be possible. Instead, studies have focused on estimating local abundance, indices of abundance, and trends of those measures using catch-per-unit-effort (CPUE), distance
(transects), and mark-recapture analyses (e.g. Epperly et al. 1995a, Leon & Diez 1999, Chaloupka & Limpus 2001, Bjorndal et al. 2005, Ehrhart et al. 2007).

The Pamlico-Albemarle Estuarine Complex of eastern North America is an important developmental and foraging habitat for loggerhead Caretta caretta, green Chelonia mydas, and Kemp’s ridley Lepidochelys kempii sea turtles (Epperly et al. 1995b). These foraging ground turtles represent several different subpopulations with origins throughout the western North Atlantic (Bass et al. 2004, 2006). Relatively large numbers of turtles are captured in pound nets set behind the barrier islands of the Complex (Epperly et al. 1995b). We present sea turtle CPUE results for these pound nets and examine trends in CPUE and size frequency distributions in order to evaluate the status of the stocks in the region.

BACKGROUND

Study area. The Pamlico-Albemarle Estuarine Complex is the largest estuarine system in the southeastern United States and is the third largest system in North America (Gross 1972). The Complex, located in North Carolina (NC), is a series of coastal lagoons, separated from the ocean by barrier islands (Fig. 1). It comprises 7 sounds: Albemarle, Currituck, Roanoke, Croatan, Pamlico, Core, and Back. The physiographic and hydrologic regimes create diverse estuarine habitats: open waters of the sounds, deeper central basins, embayments and tributary creeks, and shallow shelf areas where seagrasses abound. The area encompasses important developmental habitat for many estuarine and marine species and supports many recreational and commercial fisheries (Ross & Epperly 1985).

Pound net fisheries. The pound net was introduced in NC in the late 19th century (True 1887). Several different pound net fisheries occur in the Pamlico-Albemarle Estuarine Complex: the oldest is the late winter-early spring fishery for anadromous clupeid fishes, prosecuted well upstream, particularly in Albemarle Sound (True 1887, NCDMF 2007); a summer fishery that targets sciaenids and other marine fishes, concentrated mainly in Pamlico Sound (Higgins & Pearson 1928); and an autumn fishery that occurs in the eastern portion of the Complex, targeting southern flounder Paralichthyes lethostigma (NCDMF 2005). We worked cooperatively with the autumn flounder fishery, where pound net use is the greatest, to gain access to sea turtles.

While the size and number of nets vary by fishery, there are always 3 basic components of a pound net: a lead, a heart (named for its shape), and a pound (Fig. 2). In the case of the autumn flounder fishery, the leads are vertical walls of netting, usually 135 to 275 m in length. These leads end in the opening of the heart. The tunnel, a funnel-shaped area in the rear of the heart with a rectangular opening 70 to 90 cm and reinforced with iron rebar, connects the heart to the pound. The pounds are usually 7 to 9 m to a side, 2.4 to 4.3 m deep, and have a mesh floor. The stretch mesh size of the lead usually is 15 to 20 cm; the mesh of the heart and tunnel usually is 10 to 13 cm, and the mesh in the pound usually is 10 cm. Multiple pound nets frequently are linked together, usually in line, and collectively are named a ‘set’. Fish and turtles are intercepted by the lead and swim parallel to it, entering the heart, where they are funneled into the pound. Usually, turtles can surface in the pound to breathe, and therefore mortality is infrequent. Stakes are left in place year-round, but nets are attached only during the fishing season. Sets are registered by the State (North Carolina Marine Fisheries Commission 2007) and ownership may be transferred, but abandonment or addition of new sets occurs only rarely within a fishing season. The nets are fished by a crew of 2 to 3 persons; frequently (~20%) 2 owners fish together as a team, fishing both individuals’ nets during a trip. These
alliances usually are permanent within a season, but may vary among seasons.

Flounder fishermen begin placing their pound nets in the water during September, and continue adding nets and fishing them more frequently as the season progresses. Catches peak in November, following weather events that motivate the emigration of migratory species, including turtles. By December, flounder catch rates decline. Consequently, the nets are fished less frequently, and fishermen start to remove their nets. This fishery has existed since the early 1900s and perhaps before (N. Peele pers. comm.). Except for the introduction of gasoline outboard motors and pumps (used to put the stakes in place), this artisanal fishery has not incorporated any modern technologies.

MATERIALS AND METHODS

Our sampling occurred for 3 consecutive years (1995 to 1997), followed by 3 years with no sampling, and then 3 additional years of sampling (2001 to 2003). Flounder fishermen were chosen at random each week from the State’s registration file. Each of 4 observers attempted to sample 3 fishing trips each week, beginning mid-September (Week 1) and ending mid-December (Week 13). All nets fished by the selected fisherman and his crew that day were sampled, even if some of the sets fished belonged to a fisherman not selected (e.g. belonged to a member of the crew who also had registered nets). One observer worked with fishermen from Hatteras Island, one with fishermen from Ocracoke Island, and 2 with fishermen from Carteret County (Fig. 1), where most fishermen were based. Fishermen whose sets were located south of Oregon Inlet were selected because preliminary data indicated that turtles in the sounds to the north emigrated prior to our sampling period (NOAA Fisheries unpubl. data). Weekly aerial surveys were conducted to count sets and pounds to determine the proportion of the total fishing effort that we were sampling.

The location of each sampled pound was recorded using hand-held GPS units (Fig. 1). Calibrated thermometers were used to record surface water temperature at the first pound net fished each day. Soak time, the number of days since the pound nets were last fished, was recorded as a measure of nominal fishing effort. After turtles were brought on board, blood was drawn and tissue biopsies were taken for genetic analysis (Bass et al. 2004, 2006) and sex determination (Braun-McNeill et al. 2007). Turtles were marked (2 Inconel tags in the rear flippers and one 125 kHz PIT tag in a front flipper) to determine the actual number of individuals caught. Turtles less than 25 cm straight carapace length (SCL) were not tagged with Inconel tags. Triple tagging in this manner ensured negligible (0.005%) total mark loss (Braun-McNeill et al. 2003).

Standard straight-line and curved carapace lengths and widths were measured to identify the life stages being caught and to allow us to detect shifts in size distributions within the sampled populations. Mean sizes and size frequency distributions were compared using parametric (ANOVA) and non-parametric statistics (Kolmogorov-Smirnov, KS), respectively (SAS® Ver. 9.1 NPAR1WAY Procedure).

To estimate an annual number of individual turtles caught by species, all within-year recaptures of turtles were removed from the analysis, but recaptures between years were retained and treated as unique captures. We assumed that turtles that we were unable to mark or where we could not determine if a mark was present (e.g. escaped or too large to bring on board) were only captured once within each year, and we thus retained them in the analysis as if they were uniquely marked.

We used sets as the sampling unit in the analysis, as we could assume independence of sets, but not of pounds, which were linked within a set. To account for
the size of the sets and for the duration each pound was fished, CPUE was defined as catch per pound net-soak day and was calculated by dividing the number of captures in the sampled portion of a set by the sum of all the soak days for sampled pounds in that set. Due to the large number of zeros in the data set (sets with no captures) the delta-estimator method (Pennington 1983) was used to estimate the mean and variance of turtles captured per pound net-soak day (i.e. CPUE). Approximate 95% confidence intervals (95% CI) were calculated assuming a log-normal distribution of the CPUE. CPUEs, means, variation, coefficient of variations (CV) and CIs were calculated using MATLAB (Ver. 7.0).

Linear regression (SAS® Ver. 9.1) was used to test for a significant change in annual mean CPUEs for each species. If no trend was detected, a power analysis was conducted using TRENDS (Gerrodette 1987, 1993) to determine both our statistical power and the number of sampling intervals needed to realize a power >0.9 and thus detect a trend comparable to those reported on the nesting beaches. In the power analysis, we used the mean of the 6 annual CVs in a 2-tailed test, setting \( \alpha = 0.05 \).

RESULTS

Fishing effort usually peaked during Weeks 7 to 10, with as many as 273 sets (872 pound nets) counted south of Oregon Inlet in 1 wk (Fig. 3a). Fishing effort during 2001 to 2003 was lower than in the earlier period of 1995 to 1997, especially in 2003 when a hurricane severely impacted the area. We sampled 6 to 55% of the sets in any given week (5 to 48% of the pounds) and 14 to 30% of the sets (12 to 29% of the pounds) within a year (Fig. 3b). Pound nets per set ranged from 1 to 22, with a median of 2 and a mean of 2.9 (SD = 2.3). Mean surface water temperatures decreased over the autumn sampling period, ranging from 23.4–25.8°C in Week 1 to 6.8–15.9°C in Week 13.

We observed a total of 1303 captures of sea turtles (Table 1, Fig. 4). Of the 2167 total sets examined, 656 sets captured turtles. We captured loggerhead sea turtles in 541 sets, green turtles in 207 sets, and Kemp’s ridley turtles in 59 sets. Catches early in the sampling season were relatively low, but as the season pro-

<table>
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gressed (and water temperature dropped) the catches increased (Fig. 4). By the end of the season, catches decreased, presumably the result of autumn emigration. Throughout the course of the study, we observed 2 dead loggerhead turtles, 14 dead green turtles, and 1 dead Kemp’s ridley turtle. The catches represented a total of 910 unique captures of loggerhead turtles (Fig. 4a), 240 green turtles (Fig. 4b), and 68 Kemp’s ridley turtles (Fig. 4c, Table 1). We used our estimates of unique captures for all subsequent analyses.

Loggerhead turtles ranged in size from 41.4 to 102.5 cm SCL, with a median of 61.6 cm and a mean of 61.2 cm (SD = 7.42, n = 866) (Fig. 5). Mean size and the size distribution varied significantly among years (ANOVA, $F = 12.4532$, $p < 0.0001$, df = 5; KS = 0.1342, $p < 0.01$, n = 866). Turtles captured during the later years were larger than those captured during the earlier years (ANOVA, $F = 55.7820$, $p < 0.0001$, df = 1). The shift is most apparent in 2 size classes: 55–59 cm turtles were the dominant size class in the first couple of years, but this shifted to 60–64 cm for the last 2 yr.

Green turtles ranged in size from 20.6 to 76.7 cm SCL, with a median of 32.2 cm and a mean of 33.6 cm (SD = 10.26, n = 226) (Fig. 6). While the mean size did not vary among years (ANOVA, $F = 1.1546$, $p = 0.3329$, SCL: straight carapace length

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Fig. 4. *Caretta caretta*, *Chelonia mydas*, and *Lepidochelys kempii*. Weekly catches of (a) loggerhead, (b) green, and (c) Kemp’s ridley sea turtles, September to December 1995 to 1997 and 2001 to 2003 in eastern Pamlico, Core, and Back Sounds. Week 1 begins in mid-September (Sep 14 to 17) and Week 13 ends in mid-December (Dec 13 to 16). Note the different scales on the y-axes.

Fig. 5. *Caretta caretta*. Annual size frequency distributions of loggerhead sea turtles captured September to December in eastern Pamlico, Core, and Back Sounds, 1995 (n = 100), 1996 (n = 85), 1997 (n = 142), 2001 (n = 217), 2002 (n = 182), and 2003 (n = 140). Approximate average size of putative first time nesters (defined as unmarked turtles on beaches with saturation tagging, some of which may have been previously undetected nesters) is 90 cm (NMFS SEFSC 2001). SCL: straight carapace length.
df = 5), the size frequency distribution did (KS = 0.1391, p < 0.01, n = 226), but with no apparent pattern. Annual mean SCL ranged from 30.5 cm in 1996 (n = 26) to 36.7 cm in 2003 (n = 27).

Kemp’s ridley turtles ranged in size from 27.4 to 56.3 cm SCL, with a median of 42.2 cm and a mean of 42.3 cm (SD = 7.21, n = 67) (Fig. 7). Mean size and the size distribution varied significantly among years (ANOVA, $F = 2.8713$, p < 0.05, df = 5; KS = 0.2059, p < 0.01, n = 67), but with no apparent pattern. Annual mean SCL ranged from 38.4 cm in 2003 (n = 8) to 44.6 cm in 2002 (n = 18) (note that 1995 and 1996 are not included for this range since there were only 1 and 2 turtles measured in these years, respectively).

The mean CPUEs by year for loggerhead turtles ranged from 0.028 turtles per pound net-soak day (0.020 to 0.039 95% CI) in 1995 to 0.062 turtles per pound net-soak day (0.048 to 0.080 95% CI) in 2003 (Fig. 8a) and the CV for the annual CPUE estimates ranged from 0.11 to 0.16. The mean CPUEs for green turtles ranged from 0.006 captures per pound net-soak day (0.004 to 0.009 95% CI) in 1997 to 0.017 per pound net-soak day (0.010 to 0.027 95% CI) in 2003 (Fig. 8b) and CVs ranged from 0.18 to 0.28 (mean = 0.23). The mean CPUE for Kemp’s ridley turtles ranged from

Fig. 6. *Chelonia mydas*. Annual size frequency distributions of green sea turtles captured in eastern Pamlico, Core, and Back sounds, 1995 (n = 41), 1996 (n = 26), 1997 (n = 28), 2001 (n = 61), 2002 (n = 43), and 2003 (n = 27). Approximate average size of putative first time nesters (defined as unmarked turtles on beaches with saturation tagging, some of which may have been previously undetected nesters) is 100 cm (Zurita et al. 1993, L. M. Ehrhart unpubl. data). SCL: straight carapace length

Fig. 7. *Lepidochelys kempii*. Annual size frequency distributions of Kemp’s ridley sea turtles captured in eastern Pamlico, Core, and Back sounds, 1995 (n = 1), 1996 (n = 2), 1997 (n = 10), 2001 (n = 28), 2002 (n = 18), and 2003 (n = 8). Approximate average size of putative first time nesters (defined as unmarked turtles on beaches with saturation tagging, some of which may have been previously undetected nesters) is 60 cm (Turtle Expert Working Group 2000). SCL: straight carapace length
essentially zero ($7.72 \times 10^{-5}$; $1.15 \times 10^{-5}$ to $3.95 \times 10^{-4}$ 95% CI) in 1995 to 0.007 captures per pound net-soak day (0.004 to 0.012 95% CI) in 2002 (Fig. 8c). The CVs ranged from 0.30 to 1.00 (mean = 0.51).

We found a significant positive slope in the regression of mean annual CPUE versus time for loggerhead turtles, indicating a slow increase in loggerhead CPUE in the study area from 1995 to 2003 (CPUE = $-8.68 + 0.0044 \times \text{year}$, $R^2 = 0.99$, $p < 0.001$). The slope, or annual rate of change of the CPUE was 0.0044 turtles per pound net-soak day ($1.67 \times 10^{-4}$ SE) or 13.2%. We did not detect significant linear trends in annual CPUEs for either Kemp's ridley ($R^2 = 0.372$, $p = 0.118$) or green turtles ($R^2 = 0.015$, $p = 0.357$) over the study period.

**DISCUSSION**

Sea turtles caught on the NC foraging grounds represent many age classes of both transient and resident animals and reflect both recruitment into the neritic foraging population and permanent emigration from it (Epperly et al. 1995b, Avens et al. 2003, Sasso et al. 2006). Most were juveniles; older juveniles and adults were not well represented in our samples (Figs. 5 to 7). We would like to be able to extrapolate from trends in our CPUE estimates to trends in abundance of loggerhead sea turtle populations, but this would require making 2 major assumptions. The first is that changes in pound net catch rates reflect changes in loggerhead sea turtle abundance on the foraging grounds, implying that catch rates are proportional to abundance. Mark-recapture data often are used to estimate population sizes and thus provide an independent estimate of abundance. Unfortunately, our tag recaptures among years (11 of 863 loggerhead turtles were recaptured in a subsequent year) were too few and variable to effectively estimate population sizes or catchability coefficients or to provide an independent estimate of abundance.

Numerous studies in fisheries have demonstrated that catch rates and abundance often are not proportional (Hilborn & Walters 1992, Harley et al. 2001) and a number of factors have been reported to affect catchability, the constant that relates abundance to CPUE (Paloheimo & Dickie 1964, Rothschild 1977, Walters 2003, Maunder et al. 2006). These include variable efficiency of effort, species targeting, non-random and overlapping sampling effort, and environmental factors. As there were no changes in the fishing gear and how it was set and fished over the duration of our study, we do not think that fishing efficiency varied. Furthermore, sea turtles were not targeted by the fishery, and fishing effort was not determined by turtle catch rates (but could have been correlated with flounder catches). In addition, catchability could be related to effort, with a change in probability of capture being a function partly of the number of nets in the water. Other factors that can affect catchability may be of concern for our study. In the present study, the fixed nets were not placed randomly in the sounds, but were placed where fishing for flounder historically was productive. Thus, even though we selected the fishermen randomly and the turtles were caught passively, we did not sample the sound randomly. There is also a temporal effect that we did not address. Fishing effort and catch rates varied greatly within a season (Figs. 3a & 4), likely due to environmental factors. Early in the season, before the onset of migration, turtle catch rates generally were low, and peaked later in the season, after which they declined as the turtles left the sounds.
We pooled data within a season to calculate an annual CPUE, which means that the units of time (e.g. season) were likely not sufficiently small to assume that we effectively sampled the fishing season randomly (Walters 2003).

The second major assumption necessary to draw conclusions about trends in sea turtle populations based on our results is that the trends in CPUEs in the NC sounds represent the trends of the populations throughout their range. We are less sure about this assumption. Life history models for sea turtles, especially loggerheads, are evolving (e.g. Carr 1987, Witzell 2002, Bolten 2003). We now know that juvenile animals entering the neritic environment after an extended period in the oceanic environment do not always remain near the shore, but can shift between the 2 habitats (Keinath 1993, Witzell 2002, Mansfield 2006, McClellan & Read 2007). Furthermore, movement by adults among habitats can be size dependent (Hawkes et al. 2006, Hatase et al. 2007). Only the neritic juvenile life stages are represented in our samples. It is not known what proportion of each population enters the sounds or if that proportion is constant among years, as transiency is high (Sasso et al. 2006). In fact, the shift in size frequencies of loggerhead turtles could indicate that we sampled a different segment of the population each year.

In the absence of independent estimates of abundance to validate our CPUEs as indices of abundance, we look to other study sites along the eastern coast of North America for evidence of similar trends. Loggerhead turtles were the most frequent species captured in our study (Table 1). The NC loggerhead foraging population is dominated by turtles originating in the south Florida nesting assemblage (estimated at 80% by Bass et al. 2004). The remainder of this foraging population comprises the northern subpopulation (NC to Northern Florida) (12%), the Yucatán Peninsula (6%), and other rookeries (2%). The south Florida subpopulation was estimated to be increasing at 3.6 % yr^{-1} for the period 1981 to 1998 (Turtle Expert Working Group 2000), although in recent years the number of nests has declined (Florida Fish and Wildlife Conservation Commission 2006). The northern subpopulation, which is smaller than the south Florida assemblage by 1 order of magnitude, and as represented by surveys in South Carolina was declining at 3% yr^{-1} from 1980 to 2002 (Hopkins-Murphy et al. 2001, SCDDR 2006, S. Murphy pers. comm.). Nesting in Quintana Roo, Yucatán, was increasing 1987 to 2001 (Zurita et al. 2003), but has since decreased (J.C. Zurita pers. comm.). Thus, the NC foraging grounds represent a non-random composite of recruitment to the neritic juvenile population from all the aforementioned nesting assemblages (Bass et al. 2004, Bowen et al. 2004).

We observed a significant increase in loggerhead CPUE of 13.2% yr^{-1} (Fig. 8) during the study period, along with a significant change in the size frequencies of loggerhead turtles, represented by a shift to larger individuals over the study period. Ehrhart et al. (2007) found a similar shift in the size frequency of loggerhead turtles in the Indian River Lagoon, Florida, since the early 1990s and, while they found no significant trend in CPUE over a 24 yr period, they did report a significant increase in CPUE over the last 4 yr of their study (catch rates in 2002 to 2005 were about twice those from the earlier time periods). Sixty kilometers to the south of the Indian River Lagoon study site, at the St. Lucie Power Plant on Hutchinson Island, sea turtles are entrained with cooling water from the Atlantic Ocean into a canal system where they are net captured and released. The number of loggerhead captures at this site has increased at an average rate of more than 11 % yr^{-1} from 1988 to 2005, and a similar size shift was documented (M. Bresette pers. comm.). Lastly, shallow water trawl surveys off the southeastern coast of the USA from 1990 to 2005 also demonstrated the same size shift in loggerheads and an overall increase in sea turtle catch rates from 1990 to 2006 of about 5% yr^{-1} (SCDDR unpubl., SEAMAP-SA data). In all 3 of these other study sites, loggerheads from the south Florida subpopulation also dominated the foraging populations (Witzell et al. 2002, Roberts et al. 2005, Reece et al. 2006). CPUE trends from our study, when combined with results of these other investigations, provide evidence for an increase in the recent past in neritic juvenile loggerhead abundance in the waters of the southeastern United States.

The shift in size reported for loggerhead turtles at all of these sites could represent the progression of a dominant age class through the population, a shift in size of recruitment from the oceanic environment, sampling of different portions of the foraging population, or a reduction in recruitment. If we assume that our CPUE estimates are proportional to abundance, an increasing trend in catch rates is inconsistent with a reduction in recruitment. We note that the 2 long-term studies in Florida indicate that the average sizes in the early 2000s were comparable to those in the early 1980s, with turtles captured in the early 1990s about 4 to 5 cm smaller (Ehrhart et al. 2007, M. Bresette pers. comm.). Thus, the observation of larger turtles in recent years represents a shift to what was observed 2 decades earlier.

Green turtles were the second most abundant species captured (Table 1). Bass et al. (2006) estimated that turtles from the east coast of the United States (54%), Mexico (27%), and Costa Rica (7%) dominated the foraging population in the NC sounds. The remaining turtles likely originated in the South Atlantic.
Recent estimates indicate that the 3 dominant nesting assemblages represented are increasing: Florida at 14% yr⁻¹ (Chaloupka et al. in press), Mexico at 11% yr⁻¹ (A. Abreu pers. comm.), and Costa Rica at 5% yr⁻¹ (Troëng & Rankin 2005, Chaloupka et al. in press).

Despite an estimated linear increase greater than 10% yr⁻¹ for the 2 most dominant green turtle nesting assemblages, we did not detect a significant trend in annual CPUEs of green turtles during 1995 to 2003. However, our statistical power (0.56) may not have been sufficient to detect a trend. A power analysis indicated that 8 sampling intervals (14 yr according to our sampling design) would be needed to detect an increasing linear trend of 10% yr⁻¹. In Florida, Ehrhart et al. (2007) did find a significant positive non-linear trend in green turtle CPUE for the period 1982 to 2005 in the Indian River Lagoon, which also is dominated by turtles originating from Florida, Costa Rica, and Mexico (Bagley 2003, Bolker et al. 2007). Indian River Lagoon green turtle CPUE increased exponentially ~11% yr⁻¹ (937% overall). Results from the nearby St. Lucie Power Plant are similar: capture rates for green turtles at this site (excluding recapture events) increased from a mean of 21.1 turtles yr⁻¹ during 1977–1991 to a mean of 199.8 turtles yr⁻¹ during 1992–2006 (M. Bressette pers. comm.). Ehrhart et al. (2007) did not find any shift in size among years, although there was seasonal variability. Bjorndal et al. (2005) used mark–recapture data to examine trends in abundance in the Bahamas, where the increasing Costa Rica nesting assemblage is the major source of green turtles (Lahanas et al. 1998), and found no significant overall trend. However, during a 24 yr period at one location there were periods of significant trends (increasing followed by decreasing, and most recently stable or increasing).

Kemp’s ridley turtles were the least frequent species captured (Table 1) and CVs of the annual CPUE estimates were the highest of all 3 species. This species nests in the western Gulf of Mexico, primarily on the beaches of Tamaulipas, and nest numbers have been increasing exponentially at a rate of 14 to 16% yr⁻¹ (Heppell et al. 2005). We caught more Kemp’s ridley turtles during the latter 3 yr of our study (n = 55), particularly in 2001 and 2002, than during the first 3 yr (n = 15) (Table 1), but because of so few captures the variability in our CPUE was high and our statistical power was low (0.27). Until 2003, the catch rates were increasing (Fig. 8), but the very low catch of turtles in 2003 rendered the overall relationship insignificant. Ten sampling intervals (19 yr) would be required to detect an exponential increase of 10% yr⁻¹ on the NC foraging grounds.

To date only a few sites have been established to monitor sea turtles while in the neritic environment of eastern North America, and most of those were not established for the purpose of estimating abundance. We suggest that there is a need for long-term studies of sea turtles on foraging grounds at multiple sites throughout their ranges to estimate stock sizes and to monitor trends in CPUE and catchability as an index of their abundance. Such synoptic studies will allow us to monitor the status of multiple subpopulations of juvenile sea turtles on the foraging grounds, well before the shifts in abundance would show up on the nesting beaches. However, like nesting beach data, the high variance and the low power to detect small trends, as found for 2 species in the NC foraging ground study, indicate that projects will require either many years of sampling or an increase in sampling effort within years, or both, to decrease CVs.

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LITERATURE CITED


Bagley DA (2003) Characterizing juvenile green turtles (Chelonia mydas), from three east central Florida developmental habitats. MS thesis, University of Central Florida, Orlando, FL


Bolker BM, Okuyama T, Bjorndal KA, Bolten AB (2007) In-


SCDNR (South Carolina Department of Natural Resources) (2006) Loggerhead nesting index. SCDNR Marine Turtle Conservation Program, Charleston, SC. Available at: www.dnr.sc.gov/seaturtle/lhni.htm


